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OM protein - protein search, using sw model

Run on: January 29, 2002, 12:49:36 : Search time 38.4 Seconds
(without alignments)
192.899 Million cell updates/sec

Title: US-09-710-239-29
Perfect score: 580
Sequence: 1 RGDKGTGEGDRIKIGHRG.....DAGPVGPPGPPGPPGPP 100

Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 522463 seqs, 74073290 residues

Total number of hits satisfying chosen parameters: 522463

Minimum DB seq length: 0
Maximum DB seq length: 2000000000
Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

Database : A_Geneseq_1101.*
1: /SID88/gcgdata/geneseq/geneseq/AA1980.DAT.*
2: /SID88/gcgdata/geneseq/geneseq/AA1981.DAT.*
3: /SID88/gcgdata/geneseq/geneseq/AA1982.DAT.*
4: /SID88/gcgdata/geneseq/geneseq/AA1983.DAT.*
5: /SID88/gcgdata/geneseq/geneseq/AA1984.DAT.*
6: /SID88/gcgdata/geneseq/geneseq/AA1985.DAT.*
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9: /SID88/gcgdata/geneseq/geneseq/AA1988.DAT.*
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12: /SID88/gcgdata/geneseq/geneseq/AA1991.DAT.*
13: /SID88/gcgdata/geneseq/geneseq/AA1992.DAT.*
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17: /SID88/gcgdata/geneseq/geneseq/AA1996.DAT.*
18: /SID88/gcgdata/geneseq/geneseq/AA1997.DAT.*
19: /SID88/gcgdata/geneseq/geneseq/AA1998.DAT.*
20: /SID88/gcgdata/geneseq/geneseq/AA1999.DAT.*
21: /SID88/gcgdata/geneseq/geneseq/AA2000.DAT.*
22: /SID88/gcgdata/geneseq/geneseq/AA2001.DAT.*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	580	100.0	100	22	AAE02715
2	580	100.0	100	22	Recombinant human
3	580	100.0	200	22	Amino acid sequenc
4	580	100.0	200	22	Recombinant human
5	580	100.0	219	21	AAE02714
6	580	100.0	219	21	Amino acid sequenc
7	580	100.0	219	21	A C-terminal fragm
8	580	100.0	219	21	A C-terminal 219 ami
9	580	100.0	333	22	Recombinant human
10	580	100.0	333	22	Amino acid sequenc
11	580	100.0	441	22	Human colon cancer
	580	100.0	449	21	Human cancer assoc

12	580	100.0	510	22	AAE02712	Recombinant human
13	580	100.0	510	22	AAE02712	Amino acid sequenc
14	580	100.0	662	22	AAE02718	Human alpha (I) t
15	580	100.0	662	22	AAE02718	Amino acid sequenc
16	580	100.0	1057	21	AAE02718	Amino acid sequenc
17	580	100.0	1057	21	AAE02718	A human collagen I
18	580	100.0	1058	21	AAE02718	Amino acid sequenc
19	580	100.0	1107	17	AAE02718	Collagen/decorin(a
20	580	100.0	1107	17	AAE02718	Amino acid sequenc
21	580	100.0	1169	17	AAE02718	Collagen/BMP-28 fu
22	580	100.0	1169	17	AAE02718	Amino acid sequenc
23	580	100.0	1171	17	AAE02718	Collagen/BMP-28 fu
24	580	100.0	1171	17	AAE02718	Amino acid sequenc
25	580	100.0	1341	16	AAE02718	Collagen alpha I
26	580	100.0	1341	21	AAE02718	Collagen type I al
27	580	100.0	1388	17	AAE02718	Collagen/decorin f
28	580	100.0	1388	21	AAE02718	Amino acid sequenc
29	580	100.0	1449	22	AAE02718	Porcine alpha(I)
30	580	100.0	1464	19	AAE02718	Human recombinant
31	580	100.0	1464	22	AAE02718	Human novel protei
32	580	100.0	1464	22	AAE02718	Human pro-alpha-1
33	580	100.0	1464	22	AAE02718	Human pro-alpha-1
34	580	100.0	1464	22	AAE02718	Human pro-alpha-1
35	580	100.0	1464	22	AAE02718	Human pro-alpha-1
36	580	100.0	1464	22	AAE02718	Human pro-alpha-1
37	580	100.0	1464	22	AAE02718	Human pro-alpha-1
38	580	100.0	1464	22	AAE02718	Human pro-alpha-1
39	580	100.0	1464	22	AAE02718	Human pro-alpha-1
40	580	100.0	1464	22	AAE02718	Human pro-alpha-1
41	580	100.0	1464	22	AAE02718	Human pro-alpha-1
42	580	100.0	1464	22	AAE02718	Human pro-alpha-1
43	580	100.0	1464	22	AAE02718	Human pro-alpha-1
44	580	100.0	1464	22	AAE02718	Human pro-alpha-1
45	580	100.0	1464	22	AAE02718	Human pro-alpha-1

ALIGNMENTS

RESULT 1
AAE02715
ID AAE02715 standard; Protein; 100 AA.

XX AAE02715;

DT 06-AUG-2001 (first entry)

XX Recombinant human gelatin #4.

XX Human; recombinant gelatin; binding agent; stabilising agent; emulsifier;
encapsulant; film-forming agent; moisturising agent; thickening agent;
gelling agent; colloidal agent; adhesive agent; gel capsule; photography;
plasma expander; colloidal volume replacement material; graft coating;
medical sponge; medical plug; micro-carrier; edible composition;
protein supplement; fat substitute; nutritional supplement; cell culture;
edible coating; cosmetic; vaccine; therapy; arthritis; attheros;
cartilage degeneration; joint flexibility; food industry; beverage.

OS Homo sapiens.

XX WO200134646-A2.

PN 17-MAY-2001.

PD 10-NOV-2000; 2000WO-US30791.

XX 12-NOV-1999; 99US-0165114.

PR 15-MAY-2000; 2000US-0204437.

XX (FIBR-) FIBROGEN INC..

XX Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;

CC by that cell for naturally occurring codons not preferred by the cell;
CC incorporating the nucleic acid sequence into the cell; and contacting
CC the cell with a hypertonic growth medium containing at least one amino
CC acid, selected from the group consisting of trans-4-hydroxyproline and
CC 3-hydroxyproline to allow at least one of the amino acids to be
CC assimilated into the cell and incorporated into the extracellular matrix
CC protein. The method may be used to make host cells assimilate and
CC incorporate trans-4-hydroxyproline into proteins. This is especially
CC useful in the recombinant production of proteins such as collagen,
CC fibrinogen and fibronectin whose ability to self aggregate and produce
CC functional proteins depends on the post translational hydroxylation of
CC proline. The method is also useful in studying the structure and function
CC of polypeptides which do not normally contain trans-4-hydroxyproline.
CC The present sequence represents a C-terminal fragment of human collagen
CC type 1 (alpha1), with optimised codon usage, designated D4.
XX
XX
SQ Sequence 219 AA;

Query Match 100.0%; Score 580; DB 21; Length 219;
Best Local Similarity 100.0%; Pred. No. 3.1e-35;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTEGEGDRGIKGRGFCGSLQGPPGPGSGQPGSGAGPAGPGPGSAGAPGK 60
Db 94 rgdkgetgegdrgikgrgfsqglqgpppgpsgagpgprgpggsagapgk 153
QY 61 DGLNGLPGPIGPPGRTGDAGPVGPPGPPGPPGPPGPP 100
Db 154 dglnglpigppgrtgtdagpvppgppgppgppgpp 193

RESULT 6
AAY84555
ID AAY84555 standard; Protein; 219 AA.
XX
AC AAY84555;
XX
DT 25-JUL-2000 (first entry)
XX
DE A C-terminal fragment of human collagen type 1 (alpha2).
XX
KW Extracellular matrix protein; self aggregation; hydroxylated proline;
KW trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;
KW collagen; fibrinogen; fibronectin; post translational hydroxylation.
XX
OS Homo sapiens.
XX
PN EP92586-A2.
XX
PD 12-APR-2000.
XX
PF 07-OCT-1999; 99EP-0119184.
XX
PR 09-OCT-1998; 98US-0169768.
XX
PA (USSU) US SURGICAL CORP.
PI Gruskin EA, Buechter DD, Zhang G, Connolly K;
DR WPI; 2000-259138/23.
XX
PT Production of extracellular matrix proteins containing
PT 4-trans-hydroxyproline results in native self aggregating proteins,
PT useful on medical implants -
XX
PS Claim 10; Fig 80; 260pp; English.
XX
CC The specification describes a method for producing an extracellular
CC matrix protein or its fragment. The extracellular matrix protein is
CC capable of self aggregating in a cell which does not ordinarily
CC hydroxylated prolines. The method comprises optimising a nucleic acid
CC sequence for expression in the cell by substitution of codons preferred

CC by that cell for naturally occurring codons not preferred by the cell;
CC incorporating the nucleic acid sequence into the cell; and contacting
CC the cell with a hypertonic growth medium containing at least one amino
CC acid, selected from the group consisting of trans-4-hydroxyproline and
CC 3-hydroxyproline to allow at least one of the amino acids to be
CC assimilated into the cell and incorporated into the extracellular matrix
CC protein. The method may be used to make host cells assimilate and
CC incorporate trans-4-hydroxyproline into proteins. This is especially
CC useful in the recombinant production of proteins such as collagen,
CC fibrinogen and fibronectin whose ability to self aggregate and produce
CC functional proteins depends on the post translational hydroxylation of
CC proline. The method is also useful in studying the structure and function
CC of polypeptides which do not normally contain trans-4-hydroxyproline.
CC The present sequence represents a C-terminal fragment of human collagen
CC type 1 (alpha2), with optimised codon usage, designated D4.
XX
XX
SQ Sequence 219 AA;

Query Match 100.0%; Score 580; DB 21; Length 219;
Best Local Similarity 100.0%; Pred. No. 3.1e-35;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTEGEGDRGIKGRGFCGSLQGPPGPGSGQPGSGAGPAGPGPGSAGAPGK 60
Db 94 rgdkgetgegdrgikgrgfsqglqgpppgpsgagpgprgpggsagapgk 153
QY 61 DGLNGLPGPIGPPGRTGDAGPVGPPGPPGPPGPPGPP 100
Db 154 dglnglpigppgrtgtdagpvppgppgppgppgpp 193

RESULT 7
AAY84402
ID AAY84402 standard; Protein; 219 AA.
XX
AC AAY84402;
XX
DT 12-JUL-2000 (first entry)
XX
DE C-terminal 219 amino acids of human alpha1 collagen.
XX
KW Alpha1 collagen; 3,4-dehydro-L-proline; epoxidation; 3,4-epoxyproline;
KW collagen; mussel adhesive protein; bioadhesive.
XX
OS Homo sapiens.
XX
PN WO200014201-A1.
XX
PD 16-MAR-2000.
XX
PF 07-SEP-1999; 99WO-US20462.
XX
PR 09-SEP-1998; 98US-0099652.
XX
PA (USSU) US SURGICAL CORP.
PA (PAOL/) PAOLELLA D N.
PA (GRUS/) GRUSKIN E A.
PA (BUEC/) BUECHTER D D.
XX
PI Paolella DN, Gruskin EA, Buechter DD;
XX
DR WPI; 2000-271051/23.
DR N-PSDB; AA299842.
XX
PT Incorporating non-natural amino acid into polypeptide, useful e.g. for
PT production of bioadhesives, by epoxidation or substitution of
PT dehydroproline residues -
XX
PS Disclosure; Fig 4; 66pp; English.
XX
CC The present sequence represents the C-terminal 219 amino acids of
CC the human alpha1 collagen protein. Peptides derived from the protein

CC were used to demonstrate incorporation of 3,4-dehydro-L-proline into
CC the peptide, using the method of the invention. The specification
CC describes a method for the incorporation of non-natural amino acid
CC into a polypeptide. The method comprises reacting at least one
CC 3,4-dehydroproline residue in the polypeptide with an epoxidation
CC reagent from a polypeptide containing at least one 3,4-epoxyproline
CC residue. The method is used for studying the effects of non-natural
CC amino acids on structure and function of polypeptides. The method is
CC also useful for commercial production of collagen or mussel adhesive
CC proteins (which are useful as bioadhesives), and for incorporating a
CC wide variety of groups, including therapeutic ligands and biological
CC probes, into polypeptides.

XX
SQ Sequence 219 AA;

Query Match 100.0%; Score 580; DB 21; Length 219;
Best Local Similarity 100.0%; Pred. No. 3.1e-35;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGEGDGRGKGRGFGSLQGPDPGSGQSGAGPAGPRGPGSAGAPGK 60
|||||
DB 94 rgdkgtgeqgdrgrkgrgfgslqgppgpgsgsgagpgprgppgsagapgk 153
|||||

QY 61 DGLNGLPGPIGPPGRGTGDAGVPDPGPPGPPGPP 100
|||||
DB 154 dglnglp GPIPPGRGTGDAGVPDPGPPGPPGPP 193
|||||

RESULT 8
AAE02713
ID AAE02713 standard; Protein; 333 AA.
AC AAE02713;
XX
DT 06-AUG-2001 (first entry)
XX
DE Recombinant human gelatin #2.
DE
KW Human; recombinant gelatin; binding agent; stabilising agent; emulsifier;
KW encapsulant; film-forming agent; moisturising agent; thickening agent;
KW gelling agent; colloidal agent; adhesive agent; gel capsule; photography;
KW plasma expander; colloidal volume replacement material; graft coating;
KW medical sponge; medical plug; micro-carrier; edible composition;
KW protein supplement; fat substitute; nutritional supplement; cell culture;
KW edible coating; cosmetic; vaccine; therapy; arthritis; athrosis;
KW cartilage degeneration; joint flexibility; food industry; beverage.

OS Homo sapiens.
XX
XX WO200134646-A2.
XX
XX PD 17-MAY-2001.
XX
XX PF 10-NOV-2000; 2000WO-US30791.
XX
XX PR 12-NOV-1999; 99US-0165114.
XX
XX PR 15-MAY-2000; 2000US-0204437.
XX
XX PA (FIBR-) FIBROGEN INC.
XX
XX PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX
XX DR WPI; 2001-329072/34.
XX
XX PT Gelatin useful for pharmaceuticals, cosmetics and edible foods, is
XX prepared recombinantly -
XX
XX PS Example 1; Page 132-133; 137pp; English.
XX
XX CC The patent discloses recombinant human gelatin which is useful
XX in various compositions including binding agents, encapsulants,
XX stabilising agents, film-forming agents, moisturising agents,

CC emulsifiers, thickening agents, gelling agents, colloidal agents,
CC adhesive agents, pharmaceutical compositions, hard gel capsules,
CC soft gel capsules, plasma expander, colloidal volume replacement
CC materials, graft coatings, medical sponges, medical plugs,
CC pharmaceutical stabilisers, micro-carriers, edible compositions,
CC protein supplements, fat substitutes, nutritional supplements,
CC edible coatings, photographic compositions, cosmetic compositions,
CC industrial composition, cell culture compositions and compositions
CC for use in the laboratory. Pharmaceutical compositions comprising
CC recombinant gelatin are used as vaccines. They are also used to
CC treat various joint conditions such as arthritis, athrosis and
CC other conditions related to the degeneration of cartilage and joint
CC flexibility. Recombinant gelatin is also used in food and beverage
CC industries. The present sequence is a recombinant human gelatin.

XX
SQ Sequence 333 AA;

Query Match 100.0%; Score 580; DB 22; Length 333;
Best Local Similarity 100.0%; Pred. No. 4.3e-35;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGEGDGRGKGRGFGSLQGPDPGSGQSGAGPAGPRGPGSAGAPGK 60
|||||
DB 234 rgdkgtgeqgdrgrkgrgfgslqgppgpgsgsgagpgprgppgsagapgk 293
|||||

QY 61 DGLNGLPGPIGPPGRGTGDAGVPDPGPPGPPGPP 100
|||||
DB 294 dglnglp GPIPPGRGTGDAGVPDPGPPGPPGPP 333
|||||

RESULT 9
AAB68067
ID AAB68067 standard; Protein; 333 AA.
XX
AC AAB68067;
XX
DT 09-JUL-2001 (first entry)
XX
DE Amino acid sequence of a recombinant human gelatin.
XX
KW Human; gelatin; vaccine; anaphylactic reaction.
XX
OS Homo sapiens.
XX
XX WO200134801-A2.
XX
XX PD 17-MAY-2001.
XX
XX PF 10-NOV-2000; 2000WO-US30843.
XX
XX PR 12-NOV-1999; 99US-0165114.
XX
XX PR 15-MAY-2000; 2000US-0204437.
XX
XX PA (FIBR-) FIBROGEN INC.
XX
XX PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX
XX DR WPI; 2001-308784/32.
XX
XX PT Vaccine formulations (I) comprising recombinant human gelatin, useful
XX for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies
XX and cholera, the gelatin is non-immunogenic and confers stability at
XX ambient temperatures -
XX
XX PS Claim 11; Page 125-126; 130pp; English.
XX
XX CC The present sequence represents a human recombinant gelatin polypeptide.
XX CC The recombinant gelatin polypeptide is used to produce vaccine
XX CC formulations of the invention. The recombinant human gelatin is
XX CC non-immunogenic (therefore reducing anaphylactic reactions) and confers
XX CC stability at ambient temperatures. The vaccine formulation comprises a
XX CC vaccine formulated for the prevention of a disease selected from vaccinia

CC dermatological; neuroprotective; cardiant; thrombolytic; coagulant;
 CC neotropic; vasotropic; antipsoriatic and antiangiogenic. The
 CC polynucleotides and polypeptides can be used for preventing, treating or
 CC ameliorating medical conditions and diagnosing pathological conditions.
 CC Polynucleotides, polypeptides, antibodies, agonists and antagonists from
 CC the present invention may be used to treat immune disorders by activating
 CC or inhibiting the proliferation, differentiation or mobilisation of
 CC immune cells, to treat disorders of haematopoietic cells, autoimmune
 CC disorders, allergic reactions, graft versus host disease and organ
 CC rejection, modulate haemostatic or thrombolytic activity, modulate
 CC inflammation, cancers, cardiovascular disorders, neurological disease and
 CC bacterial or viral infections. The peptides, nucleotides, antibodies,
 CC agonists and antagonists may be also be used in drug screens. AAC78449 to
 CC AAC78457 and AAB44240 represent sequences used in the exemplification of
 CC the present invention.

XX Sequence 449 AA;

Query Match 100.0%; Score 580; DB 21; Length 449;
 Best Local Similarity 100.0%; Pred. No. 5.4e-35;
 Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RDKGETGEQDGRGKIHGRFSGLGQPPGPGSPGEGQPGSGAGPAGPPGSGAGPCK 60
 Db 78 rgdkgetgegdrgikghrgfsglgqpppgpsgeqpgsgagpagprgpggsagapgk 137
 QY 61 DGLNGLPGPIGPPGPRGRTGDAGVPVPGPPGPPGPPGPPGPP 100
 Db 138 dglnglpigppgprgrtgdagvpvpgpppgpppgpppp 177

RESULT 12

AAE02712
 ID AAE02712 standard; Protein; 510 AA.

XX AAE02712;

DT 06-AUG-2001 (first entry)

DE Recombinant human gelatin #1.

XX Human; recombinant gelatin; binding agent; stabilising agent; emulsifier;
 KW encapsulant; film-forming agent; moisturising agent; thickening agent;
 KW gelling agent; colloidal agent; adhesive agent; gel capsule; photography;
 KW plasma expander; colloidal volume replacement material; graft coating;
 KW medical sponge; medical plug; micro-carrier; edible composition;
 KW protein supplement; fat substitute; nutritional supplement; cell culture;
 KW edible coating; cosmetic; vaccine; therapy; arthritis; athrosis;
 KW cartilage degeneration; joint flexibility; food industry; beverage.

XX Homo sapiens.

OS WO200134646-A2.

PN 17-MAY-2001.

XX 10-NOV-2000; 2000WO-US30791.

PR 12-NOV-1999; 99US-0165114.

PR 15-MAY-2000; 2000US-0204437.

XX (FIBR-) FIBROGEN INC.

PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;

XX WPI; 2001-329072/34.

XX Gelatin useful for pharmaceuticals, cosmetics and edible foods, is
 PT prepared recombinantly -

XX Disclosure; Page 130-131; 137pp; English.

CC The patent discloses recombinant human gelatin which is useful
 CC in various compositions including binding agents, encapsulants,
 CC stabilising agents, film-forming agents, moisturising agents,
 CC emulsifiers, thickening agents, gelling agents, colloidal agents,
 CC adhesive agents, pharmaceutical compositions, hard gel capsules,
 CC soft gel capsules, plasma expander, colloidal volume replacement
 CC materials, graft coatings, medical sponges, medical plugs,
 CC pharmaceutical stabilisers, micro-carriers, edible compositions,
 CC protein supplements, fat substitutes, nutritional supplements,
 CC edible coatings, photographic compositions, cosmetic compositions,
 CC industrial composition, cell culture compositions and compositions
 CC for use in the laboratory. Pharmaceutical compositions comprising
 CC recombinant gelatin are used as vaccines. They are also used to
 CC treat various joint conditions such as arthritis, athrosis and
 CC other conditions related to the degeneration of cartilage and joint
 CC flexibility. Recombinant gelatin is also used in food and beverage
 CC industries. The present sequence is a recombinant human gelatin.

XX Sequence 510 AA;

Query Match 100.0%; Score 580; DB 22; Length 510;
 Best Local Similarity 100.0%; Pred. No. 6e-35;
 Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RDKGETGEQDGRGKIHGRFSGLGQPPGPGSPGEGQPGSGAGPAGPPGSGAGPCK 60
 Db 411 rgdkgetgegdrgikghrgfsglgqpppgpsgeqpgsgagpagprgpggsagapgk 470
 QY 61 DGLNGLPGPIGPPGPRGRTGDAGVPVPGPPGPPGPPGPPGPP 100
 Db 471 dglnglpigppgprgrtgdagvpvpgpppgpppgpppp 510

RESULT 13

AAB68066

ID AAB68066 standard; Protein; 510 AA.

XX AAB68066;

DT 09-JUL-2001 (first entry)

XX Amino acid sequence of a recombinant human gelatin.

XX Human; gelatin; vaccine; anaphylactic reaction.

XX Homo sapiens.

XX WO200134801-A2.

XX 17-MAY-2001.

XX 10-NOV-2000; 2000WO-US30843.

XX 12-NOV-1999; 99US-0165114.

XX 15-MAY-2000; 2000US-0204437.

XX (FIBR-) FIBROGEN INC.

XX Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;

XX WPI; 2001-308784/32.

XX Vaccine formulations (I) comprising recombinant human gelatin, useful
 PT for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies
 PT and cholera, the gelatin is non-immunogenic and confers stability at
 PT ambient temperatures -

PS Claim 11; Page 123-124; 130pp; English.

XX The present sequence represents a human recombinant gelatin polypeptide.
 CC The recombinant gelatin polypeptide is used to produce vaccine
 CC formulations of the invention. The recombinant human gelatin is

CC non-immunogenic (therefore reducing anaphylactic reactions) and confers
 CC stability at ambient temperatures. The vaccine formulation comprises a
 CC vaccine formulated for the prevention of a disease selected from vaccinia
 CC virus (small pox), polio virus (Salk and Sabin), mumps, measles, rubella,
 CC diphtheria, tetanus, Varicella-Zoster (chicken pox/shingles), pertussis
 CC (whooping cough), Bacille Calmette-Guérin (BCG, tuberculosis),
 CC haemophilus influenzae meningitis, rabies, cholera, Japanese
 CC encephalitis virus, salmonella typhi, shigella, hepatitis A and B,
 CC adenovirus, yellow fever, foot and mouth disease, herpes simplex virus,
 CC respiratory syncytial virus, rotavirus, Dengue, West Nile virus, turkey
 CC herpes virus (Marek's disease), influenza and/or anthrax.
 XX
 SQ Sequence 510 AA;

Query Match 100.0%; Score 580; DB 22; Length 510;
 Best Local Similarity 100.0%; Pred. No. 6e-35;
 Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGGDRGKIGHRGFSGLQPPGPPGSGGSGAGPAGPAGPAGPAGK 60
 Db 411 rgdkgtgeqgdrigkhrfgslgppgppgsggsgagpgrgppgsagapgk 470

QY 61 DGLNGLPGPIGPPGPRGTGACPVGPPGPPGPPGPP 100

Db 471 dglnglpgpigrtpgrtgagvpgppgppgppgpp 510

RESULT 14

AAE02718

ID AAE02718 standard; Protein; 662 AA.

XX

AC AAE02718;

XX

DT 06-AUG-2001 (first entry)

XX

DE Human alpha (I) type I collagen helical domain (residues 531-1192).

XX

KW Human; recombinant gelatin; binding agent; stabilising agent; emulsifier;
 KW encapsulant; film-forming agent; moisturing agent; thickening agent;
 KW gelling agent; colloidal agent; adhesive agent; gel capsule; photography;
 KW plasma expander; colloidal volume replacement material; graft coating;
 KW medical sponge; medical plug; micro-carrier; edible composition;
 KW protein supplement; fat substitute; nutritional supplement; cell culture;
 KW edible coating; cosmetic; vaccine; therapy; arthritis; atrophosis;
 KW cartilage degeneration; joint flexibility; food industry; beverage;
 KW alpha (I) type I collagen.

XX

OS Homo sapiens.

XX

FT WO200134646-A2.

XX

PN 17-MAY-2001.

XX

PD 10-NOV-2000; 2000WO-US30791.

XX

PF 12-NOV-1999; 99US-0165114.

XX

PR 15-MAY-2000; 2000US-0204437.

XX

PA (FIBR-) FIBROGEN INC.

XX

PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;

XX

DR WPI; 2001-329072/34.

XX

PT Gelatin useful for pharmaceuticals, cosmetics and edible foods, is
 prepared recombinantly.

XX

PS Claim 21; Page 135-137; 137pp; English.

XX

CC The patent discloses recombinant human gelatin which is useful
 CC in various compositions including binding agents, encapsulants,
 CC stabilising agents, film-forming agents, moisturising agents,

XX

PS

XX

PS

XX

PS

XX

CC emulsifiers, thickening agents, gelling agents, colloidal agents,
 CC adhesive agents, pharmaceutical compositions, hard gel capsules,
 CC soft gel capsules, plasma expander, colloidal volume replacement
 CC materials, graft coatings, medical sponges, medical plugs,
 CC pharmaceutical stabilisers, micro-carriers, edible compositions,
 CC protein supplements, fat substitutes, nutritional supplements,
 CC edible coatings, photographic compositions, cosmetic compositions,
 CC industrial composition, cell culture compositions and compositions
 CC for use in the laboratory. Pharmaceutical compositions comprising
 CC recombinant gelatin are used as vaccines. They are also used to
 CC treat various joint conditions such as arthritis, atrophosis and
 CC other conditions related to the degeneration of cartilage and joint
 CC flexibility. Recombinant gelatin is also used in food and beverage
 CC industries. The present sequence is human alpha (I) type I collagen
 CC helical domain (residues 531-1192). This sequence is a recombinant
 CC gelatin.

XX
 SQ Sequence 662 AA;

Query Match 100.0%; Score 580; DB 22; Length 662;
 Best Local Similarity 100.0%; Pred. No. 7.4e-35;
 Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGGDRGKIGHRGFSGLQPPGPPGSGGSGAGPAGPAGPAGK 60
 Db 563 rgdkgtgeqgdrigkhrfgslgppgppgsggsgagpgrgppgsagapgk 622

QY 61 DGLNGLPGPIGPPGPRGTGACPVGPPGPPGPPGPP 100

Db 623 dglnglpgpigrtpgrtgagvpgppgppgppgppgpp 662

RESULT 15

AAB68072

ID AAB68072 standard; Protein; 662 AA.

XX

AC AAB68072;

XX

DT 09-JUL-2001 (first entry)

XX

DE Amino acid sequence of a recombinant human gelatin.

XX

KW Human; gelatin; vaccine; anaphylactic reaction.

XX

OS Homo sapiens.

XX

FH Key Location/Qualifiers

FT

FT Misc-difference 53 /note= "this residue is given as unknown as it is
 FT illegible in the specification"

FT

PN WO200134801-A2.

XX

PD 17-MAY-2001.

XX

PF 10-NOV-2000; 2000WO-US30843.

XX

PR 12-NOV-1999; 99US-0165114.

XX

PR 15-MAY-2000; 2000US-0204437.

XX

PA (FIBR-) FIBROGEN INC.

XX

PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;

XX

DR WPI; 2001-308784/32.

XX

PT Vaccine formulations (I) comprising recombinant human gelatin, useful
 PT for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies
 PT and cholera, the gelatin is non-immunogenic and confers stability at
 PT ambient temperatures.

XX

PS Claim 11; Page 128-130; 130pp; English.

XX

XX The present sequence represents a human recombinant gelatin polypeptide.
 CC The recombinant gelatin polypeptide is used to produce vaccine
 CC formulations of the invention. The recombinant human gelatin is
 CC non-immunogenic (therefore reducing anaphylactic reactions) and confers
 CC stability at ambient temperatures. The vaccine formulation comprises a
 CC vaccine formulated for the prevention of a disease selected from vaccinia
 CC virus (small pox), polio virus (Salk and Sabin), mumps, measles, rubella,
 CC diphtheria, tetanus, Varicella-Zoster (chicken pox/shingles), pertussis
 CC (whooping cough), Bacille Calmette-Guérin (BCG, tuberculosis),
 CC haemophilus influenzae meningitis, rabies, cholera, Japanese
 CC encephalitis virus, salmonella typhi, shigella, hepatitis A and B,
 CC adenovirus, yellow fever, foot and mouth disease, herpes simplex virus,
 CC respiratory syncytial virus, rotavirus, Dengue, West Nile virus, turkey
 CC herpes virus (Marek's disease), influenza and/or anthrax.

XX Sequence 662 AA;

Query Match 100.0%; Score 580; DB 22; Length 662;
 Best Local Similarity 100.0%; Pred. No. 7.4e-35;
 Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 RCDKGTGEQDGRGKIGHRGFSGLOGPPGPGSPGEGQPGSGAGPAGRPPGSGAGPK 60
 Db 563 rdkgetgegdrglkghrgfsgldgppppgpgsggpgsgagpagrppgsagapgk 622
 QY 61 DGLNGLPGPIGPPGPRGRTGDAGVGPVGPVGPVGPVGPVGPVGPVGPVGPVGP 100
 Db 623 dglnglpigppgprgrtgdagvvpvpppppppppppppp 662

RESULT 16

AA84541
 ID AAY84541 standard; Protein; 1057 AA.

AC AAY84541;

DT 25-JUL-2000 (first entry)

DE Amino acid sequence of a human collagen 1 (alpha1) protein.

XX Extracellular matrix protein; self aggregation; hydroxylated proline;
 KW trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;
 KW collagen; fibrinogen; fibronectin; post translational hydroxylation.

OS Homo sapiens.

PN EP992586-A2.

PD 12-APR-2000.

PF 07-OCT-1999; 99EP-0119184.

XX 09-OCT-1998; 98US-0169768.

PR (USSU) US SURGICAL CORP.

PA Gruskin EA, Buechter DD, Zhang G, Connolly K;

PI WPI; 2000-259138/23.

DR N-PSDB; AAA12502.

XX Production of extracellular matrix proteins containing
 PT 4-trans-hydroxyproline results in native self aggregating proteins,
 PT useful on medical implants -

XX Disclosure; Fig 27A-E; 260pp; English.

XX The specification describes a method for producing an extracellular
 CC matrix protein or its fragment. The extracellular matrix protein is
 CC capable of self aggregating in a cell which does not ordinarily
 CC hydroxylated prolines. The method comprises optimising a nucleic acid

CC sequence for expression in the cell by substitution of codons preferred
 CC by that cell for naturally occurring codons not preferred by the cell;
 CC incorporating the nucleic acid sequence into the cell; and contacting
 CC the cell with a hypertonic growth medium containing at least one amino
 CC acid, selected from the group consisting of trans-4-hydroxyproline and
 CC 3-hydroxyproline to allow at least one of the amino acids to be
 CC assimilated into the cell and incorporated into the extracellular matrix
 CC protein. The method may be used to make host cells assimilate and
 CC incorporate trans-4-hydroxyproline into proteins. This is especially
 CC useful in the recombinant production of proteins such as collagen,
 CC fibrinogen and fibronectin whose ability to self aggregate and produce
 CC functional proteins depends on the post translational hydroxylation of
 CC proline. The method is also useful in studying the structure and function
 CC of polypeptides which do not normally contain trans-4-hydroxyproline.
 CC The present sequence represents a human collagen 1 (alpha1) protein,
 CC which may be produced using the method of the invention.

XX Sequence 1057 AA;

Query Match 100.0%; Score 580; DB 21; Length 1057;
 Best Local Similarity 100.0%; Pred. No. 1.1e-34;
 Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 RCDKGTGEQDGRGKIGHRGFSGLOGPPGPGSPGEGQPGSGAGPAGRPPGSGAGPK 60
 Db 932 rdkgetgegdrglkghrgfsgldgppppgpgsggpgsgagpagrppgsagapgk 991
 QY 61 DGLNGLPGPIGPPGPRGRTGDAGVGPVGPVGPVGPVGPVGPVGPVGPVGPVGP 100
 Db 992 dglnglpigppgprgrtgdagvvpvpppppppppppppp 1031

RESULT 17

AA84544
 ID AAY84544 standard; Protein; 1057 AA.

AC AAY84544;

DT 25-JUL-2000 (first entry)

DE A human collagen 1 (alpha1) protein helical region.

XX Extracellular matrix protein; self aggregation; hydroxylated proline;
 KW trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;
 KW collagen; fibrinogen; fibronectin; post translational hydroxylation.

OS Homo sapiens.

PN EP992586-A2.

PD 12-APR-2000.

PF 07-OCT-1999; 99EP-0119184.

XX 09-OCT-1998; 98US-0169768.

PR (USSU) US SURGICAL CORP.

PA Gruskin EA, Buechter DD, Zhang G, Connolly K;

PI WPI; 2000-259138/23.

DR N-PSDB; AAA12503.

XX Production of extracellular matrix proteins containing
 PT 4-trans-hydroxyproline results in native self aggregating proteins,
 PT useful on medical implants -

XX Example 10; Fig 39A-E; 260pp; English.

XX The specification describes a method for producing an extracellular
 CC matrix protein or its fragment. The extracellular matrix protein is
 CC capable of self aggregating in a cell which does not ordinarily

CC hydroxylated prolines. The method comprises optimising a nucleic acid
 CC sequence for expression in the cell by substitution of codons preferred
 CC by that cell for naturally occurring codons not preferred by the cell;
 CC incorporating the nucleic acid sequence into the cell; and contacting
 CC the cell with a hypertonic growth medium containing at least one amino
 CC acid, selected from the group consisting of trans-4-hydroxyproline and
 CC 3-hydroxyproline to allow at least one of the amino acids to be
 CC assimilated into the cell and incorporated into the extracellular matrix
 CC protein. The method may be used to make host cells assimilate and
 CC incorporate trans-4-hydroxyproline into proteins. This is especially
 CC useful in the recombinant production of proteins such as collagen,
 CC fibrinogen and fibronectin whose ability to self aggregate and produce
 CC functional proteins depends on the post translational hydroxylation of
 CC proline. The method is also useful in studying the structure and function
 CC of polypeptides which do not normally contain trans-4-hydroxyproline.
 CC The present sequence represents human collagen 1 (alpha1) helical region,
 CC which may be produced using the method of the invention.

XX Sequence 1057 AA;

Query Match 100.0%; Score 580; DB 21; Length 1057;

Best Local Similarity 100.0%; Pred. No. 1.1e-34;

Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGEGDRGKIGHRGFSGLQGPDPGSGSGAGPAGPRGPGSAGAPGK 60

|||||
 Db 932 rgdkgtgegdrgikghrgfsglqgpppgsgsgagsgagprgppgsagapgk 991

QY 61 DGLNGLPGIPGPPGRGTGDAGPVGPPGPPGPPGPP 100

|||||
 Db 992 dginglpgipgppgrgtgdagpvpgpppppppppp 1031

RESULT 18

AAV84403

ID AAY84403 standard; Protein; 1058 AA.

XX AC AAY84403;

XX DT 12-JUL-2000 (first entry)

XX DE Amino acid sequence of human type 1 (alpha1) collagen polypeptide.

XX KW Alpha1 collagen; 3,4-dehydro-L-proline; epoxidation; 3,4-epoxyproline;

XX KW collagen; mussel adhesive protein; bioadhesive.

XX OS Homo sapiens.

XX PN WO200014201-A1.

XX PD 16-MAR-2000.

XX PF 07-SEP-1999; 99WO-US20462.

XX PR 09-SEP-1998; 98US-0099652.

XX PA (USSU) US SURGICAL CORP.

XX PA (PAOL/) PAOLELIA D N.

XX PA (GRUS/) GRUSKIN E A.

XX PA (BUEC/) BUECHTER D D.

XX PI Paolella DN, Gruskin EA, Buechter DD;

XX DR WPI; 2000-271051/23.

XX DR N-PSDB; AA299843.

XX PT Incorporating non-natural amino acid into polypeptide, useful e.g. for

XX PT production of bioadhesives, by epoxidation or substitution of

XX PT dehydroproline residues

XX PS Disclosure; Fig 6; 66pp; English.

XX

CC The present sequence represents a human type 1 (alpha1) collagen protein.
 CC Peptides derived from the protein were used to demonstrate incorporation
 CC of 3,4-dehydro-L-proline into the peptide, using the method of the
 CC invention. The specification describes a method for the incorporation
 CC of non-natural amino acid into a polypeptide. The method comprises
 CC reacting at least one 3,4-dehydroproline residue in the polypeptide
 CC with an epoxidation reagent from a polypeptide containing at least
 CC one 3,4-epoxyproline residue. The method is used for studying the
 CC effects of non-natural amino acids on structure and function of
 CC polypeptides. The method is also useful for commercial production of
 CC collagen or mussel adhesive proteins (which are useful as bioadhesives),
 CC and for incorporating a wide variety of groups, including therapeutic
 CC ligands and biological probes, into polypeptides.

XX Sequence 1058 AA;

Query Match 100.0%; Score 580; DB 21; Length 1058;

Best Local Similarity 100.0%; Pred. No. 1.1e-34;

Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGEGDRGKIGHRGFSGLQGPDPGSGSGAGPAGPRGPGSAGAPGK 60

|||||
 Db 933 rgdkgtgegdrgikghrgfsglqgpppgsgsgagsgagprgppgsagapgk 992

QY 61 DGLNGLPGIPGPPGRGTGDAGPVGPPGPPGPPGPP 100

|||||
 Db 993 dginglpgipgppgrgtgdagpvpgpppppppppp 1032

RESULT 19

AAAR89472

ID AAR89472 standard; Protein; 1107 AA.

XX AC AAR89472;

XX DT 01-OCT-1996 (first entry)

XX DE Collagen/Decorin(aa46-93) fusion protein.

XX KW Transforming growth factor; TGF-beta-1; collagen IA; osteogenesis;

XX KW bone formation; tissue repair; fusion protein.

XX OS Synthetic.

XX FH Key Location/Qualifiers

XX FT Domain 1..1057

XX FT /label= Collagen-IA

XX FT /note= "collagen IA alpha-helical domain"

XX FT Peptide 1058..1059

XX FT /label= Linker_peptide

XX FT Domain 1060..1107

XX FT /label= Decorin

XX FT /note= "amino acids P46 to G93 of mature

XX FT decorin"

XX FT Misc-difference 887

XX FT /note= "unidentified amino acid"

XX FT Misc-difference 890

XX FT /note= "unidentified amino acid"

XX PN CA2151547-A.

XX PD 11-DEC-1995.

XX PF 12-JUN-1995; 95CA-2151547.

XX PR 10-JUN-1994; 94US-0259263.

XX PA (USSU) US SURGICAL CORP.

XX PI Espino P, Gruskin EA;

XX DR WPI; 1996-140144/15.

DR N-PSDB; AAT16518.

XX Chimaeric DNA encoding protein contg. extracellular matrix protein

PT domain - and cellular regulatory factor domain, partic. useful as

PT osteogenic agents, also related vectors, transformed cells and

PT chimaeric proteins.

PS Disclosure; Fig 8; 59pp; English.

XX A fusion protein (AAR89472) comprises the alpha-helical region of

CC human collagen I(a) linked to amino acids 46-93 of human mature

CC dermatan sulphate proteoglycan (decorin). It can be expressed in

CC Escherichia coli transformants carrying a vector incorporating a

CC chimeric gene (AAT16518) coding for the fusion. The decorin binds to

CC type I collagen and thus affects Elbril formation. It inhibits

CC the cell attachment-promoting activity of collagen and fibrinogen

CC by binding to such molecules near their cell binding sites. The

CC collagen moiety provides an integral substratum or scaffolding for

CC the decorin. The fusion protein acts to reduce scarring of healing

CC tissue.

XX Sequence 1107 AA;

SQ

Query Match 100.0%; Score 580; DB 17; Length 1107;

Best Local Similarity 100.0%; Pred. No. 1.1e-34;

Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 RDKGTCGEQDGRGKIGHRGFSGLGPPGPGSPGCEQPGSGAGPGRGPGSGAGPCK 60

Db 932 rgdkgetgeqgdrglkghrgfsglqgpppgpsgeqpgsgagpagprgpggsagapgk 991

Qy 61 DGLNGLPGPIGPPGRGRTGDAGPVGPPGPGPPGPPGPP 100

Db 992 dglnlpgpippgprgrtgdagpvpgpppgpppppppp 1031

RESULT 20

AAY84540

ID AAY84540 standard; Protein; 1107 AA.

XX AAY84540;

AC AAY84540;

XX

DT 25-JUL-2000 (first entry)

XX

DE Amino acid sequence of a chimeric collagen 1 (alpha1)/decorin protein.

XX

XX Extracellular matrix protein; self aggregation; hydroxylated proline;

KW trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;

KW collagen; fibrinogen; fibronectin; post translational hydroxylation;

KW decorin; chimera.

XX

OS Chimeric - Homo sapiens.

OS Chimeric - Unidentified.

XX

FH Key Location/Qualifiers

FT Misc-difference 858

FT /note= "Gly encoded by GCT"

XX

PN EP992586-A2.

XX

XX 12-APR-2000.

XX

PF 07-OCT-1999; 99EP-0119184.

XX

XX 09-OCT-1998; 98US-0169768.

PR (USSU) US SURGICAL CORP.

PA

XX Gruskin EA, Buechter DD, Zhang G, Connolly K;

PI

XX WPI; 2000-259138/23.

DR

DR N-PSDB; AAA12500.

XX Production of extracellular matrix proteins containing

PT 4-trans-hydroxyproline results in native self aggregating proteins,

PT useful on medical implants -

XX

PS Claim 24; Fig 18; 260pp; English.

XX

CC The specification describes a method for producing an extracellular

CC matrix protein or its fragment. The extracellular matrix protein is

CC capable of self aggregating in a cell which does not ordinarily

CC hydroxylated prolines. The method comprises optimising a nucleic acid

CC sequence for expression in the cell by substitution of codons preferred

CC by that cell for naturally occurring codons not preferred by the cell;

CC incorporating the nucleic acid sequence into the cell; and contacting

CC the cell with a hypertonic growth medium containing at least one amino

CC acid, selected from the group consisting of trans-4-hydroxyproline and

CC 3-hydroxyproline to allow at least one of the amino acids to be

CC assimilated into the cell and incorporated into the extracellular matrix

CC protein. The method may be used to make host cells assimilate and

CC incorporate trans-4-hydroxyproline into proteins. This is especially

CC useful in the recombinant production of proteins such as collagen,

CC fibrinogen and fibronectin whose ability to self aggregate and produce

CC functional proteins depends on the post translational hydroxylation of

CC proline. The method is also useful in studying the structure and function

CC of polypeptides which do not normally contain trans-4-hydroxyproline.

CC The present sequence represents a chimeric collagen 1 (alpha1)/decorin

CC protein, which may be produced using the method of the invention.

XX

SQ Sequence 1107 AA;

Query Match 100.0%; Score 580; DB 21; Length 1107;

Best Local Similarity 100.0%; Pred. No. 1.1e-34;

Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 RDKGTCGEQDGRGKIGHRGFSGLGPPGPGSPGCEQPGSGAGPGRGPGSGAGPCK 60

Db 932 rgdkgetgeqgdrglkghrgfsglqgpppgpsgeqpgsgagpagprgpggsagapgk 991

Qy 61 DGLNGLPGPIGPPGRGRTGDAGPVGPPGPPGPPGPPGPP 100

Db 992 dglnlpgpippgprgrtgdagpvpgpppgpppppppp 1031

RESULT 21

AAR89469

ID AAR89469 standard; Protein; 1169 AA.

XX

AC AAR89469;

XX

DT 01-OCT-1996 (first entry)

XX

XX Collagen/BMP-2B fusion protein.

DE

XX Bone morphogenic protein 2B; BMP-2B; collagen IA; osteogenesis;

KW fusion protein.

KW

XX Synthetic.

OS

XX

FH Key Location/Qualifiers

FT Domain 1..1057

FT /label= Collagen-IA

FT /note= "collagen IA alpha-helical domain"

FT Peptide 1058..1059

FT /label= linker_peptide

FT Domain 1060..1169

FT /label= BMP-2B

FT /note= "human mature BMP-2B"

FT Misc-difference 887

FT /note= "unidentified amino acid"

FT Misc-difference 890

FT /note= "unidentified amino acid"

XX

PN CA2151547-A.
 XX
 PD 11-DEC-1995.
 XX
 PF 12-JUN-1995; 95CA-2151547.
 XX
 PR 10-JUN-1994; 94US-0259263.
 XX
 PA (USSU) US SURGICAL CORP.
 XX
 PI Espino P, Gruskin EA;
 XX
 DR WPI; 1996-140144/15.
 DR N-PSDB; AAT16515.
 XX
 PT Chimaeric DNA encoding protein contg. extracellular matrix protein
 PT domain - and cellular regulatory factor domain, partic. useful as
 PT osteogenic agents, also related vectors, transformed cells and
 PT chimaeric proteins.
 XX
 PS Disclosure; Flg 5; 59pp; English.
 XX
 CC A fusion protein (AAR89469) comprises the alpha-helical region of
 CC human collagen I(a) linked to the human mature bone morphogenic
 CC protein 2B (BMP2B). It can be expressed in Escherichia coli
 CC transformants carrying a vector incorporating a chimeric gene
 CC (AAT16515) coding for the fusion. The BMP moiety induces
 CC osteogenesis, while the collagen moiety provides an integral
 CC substratum or scaffolding for the BMP and cells involved in
 CC reconstruction and growth. The fusion protein provides sustained
 CC release and delivery of BMP to a target tissue.
 XX
 SQ Sequence 1169 AA;
 Query Match 100.0%; Score 580; DB 17; Length 1169;
 Best Local Similarity 100.0%; Pred. No. 1.2e-34;
 Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 RGDKGETGEGDGRGKGRHGFSGLOGPPGPPSPGQGPAGSGAGPAGPPGSGAGAPGK 60
 Db 932 rgdkgetgegdgrgkgrhgfsglqppppspgqgpasgagpgrgppgsagapgk 991
 QY 61 DGLNGLPGPIGPPGPRGRTGDAGVGPDPGPPGPPGPP 100
 Db 992 dglnglpgpigrgrtgadgvgppgppgppgppgpp 1031
 RESULT 22
 AAY84537
 ID AAY84537 standard; Protein; 1169 AA.
 XX
 AC AAY84537;
 XX
 DT 25-JUL-2000 (first entry)
 XX
 DE Amino acid sequence of a chimeric collagen 1 (alpha1)/BMP-2B protein.
 XX
 KW Extracellular matrix protein; self aggregation; hydroxylated proline;
 KW trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;
 KW collagen; fibrinogen; fibronectin; post translational hydroxylation; ss.
 KW bone morphogenic protein; BMP-2B; chimera.
 XX
 OS Chimeric - Homo sapiens.
 OS Chimeric - Unidentified.
 XX
 FH Key Location/Qualifiers
 FT Misc-difference 677 /note= "Ala encoded by G"
 FT Misc-difference 887 /note= "unspecified amino acid encoded by CT"
 FT Misc-difference 890 /note= "unspecified amino acid encoded by CT"
 FT

XX EP992586-A2.
 PN 12-APR-2000.
 XX
 PD 07-OCT-1999; 99EP-0119184.
 XX
 PF 09-OCT-1998; 98US-0169768.
 XX
 PR (USSU) US SURGICAL CORP.
 XX
 PA Gruskin EA, Buechter DD, Zhang G, Connolly K;
 XX
 PI WPI; 2000-259138/23.
 DR N-PSDB; AAA12497.
 XX
 PT Production of extracellular matrix proteins containing
 PT 4-trans-hydroxyproline results in native self aggregating proteins,
 PT useful on medical implants -
 XX
 PS Claim 22; Fig 13; 260pp; English.
 XX
 CC The specification describes a method for producing an extracellular
 CC matrix protein or its fragment. The extracellular matrix protein is
 CC capable of self aggregating in a cell which does not ordinarily
 CC hydroxylated prolines. The method comprises optimising a nucleic acid
 CC sequence for expression in the cell by substitution of codons preferred
 CC by that cell for naturally occurring codons not preferred by the cell;
 CC incorporating the nucleic acid sequence into the cell; and contacting
 CC the cell with a hypertonic growth medium containing at least one amino
 CC acid, selected from the group consisting of trans-4-hydroxyproline and
 CC 3-hydroxyproline to allow at least one of the amino acids to be
 CC assimilated into the cell and incorporated into the extracellular matrix
 CC protein. The method may be used to make host cells assimilate and
 CC incorporate trans-4-hydroxyproline into proteins. This is especially
 CC useful in the recombinant production of proteins such as collagen,
 CC fibrinogen and fibronectin whose ability to self aggregate and produce
 CC functional proteins depends on the post translational hydroxylation of
 CC proline. The method is also useful in studying the structure and function
 CC of polypeptides which do not normally contain trans-4-hydroxyproline.
 CC The present sequence represents a chimeric collagen 1 (alpha1)/bone
 CC morphogenic protein-2B (bmp-2b) protein, which may be produced using the
 CC method of the invention.
 XX
 SQ Sequence 1169 AA;
 Query Match 100.0%; Score 580; DB 21; Length 1169;
 Best Local Similarity 100.0%; Pred. No. 1.2e-34;
 Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 RGDKGETGEGDGRGKGRHGFSGLOGPPGPPSPGQGPAGSGAGPAGPPGSGAGAPGK 60
 Db 932 rgdkgetgegdgrgkgrhgfsglqppppspgqgpasgagpgrgppgsagapgk 991
 QY 61 DGLNGLPGPIGPPGPRGRTGDAGVGPDPGPPGPPGPP 100
 Db 992 dglnglpgpigrgrtgadgvgppgppgppgppgpp 1031
 RESULT 23
 AAR89470
 ID AAR89470 standard; Protein; 1171 AA.
 XX
 AC AAR89470;
 XX
 DT 01-OCT-1996 (first entry)
 XX
 DE Collagen/TGF-beta-1 fusion protein.
 XX
 KW Transforming growth factor; TGF-beta-1; collagen 1A; osteogenesis;
 KW bone formation; tissue repair; fusion protein.
 XX

```
OS Synthetic.
XX Key Location/Qualifiers
FH 1..1057
FT /label= Collagen-1A
FT /note= "collagen 1A alpha-helical domain"
FT 1058..1059
FT /label= Linker_peptide
FT 1060..1171
FT /label= TGF-beta-1
FT /note= "human mature TGF-beta-1"
FT Misc-difference 887
FT /note= "unidentified amino acid"
FT Misc-difference 890
FT /note= "unidentified amino acid"
XX CA21513177A.
XX
XX PD 11-DEC-1995.
XX
XX PD 12-JUN-1995; 95CA-2151547.
XX
XX PR 10-JUN-1994; 94US-0259263.
XX
XX PA (USSU ) US SURGICAL CORP.
XX
XX PI Espino P, Gruskin EA;
XX
XX DR WPI; 1996-140144/15.
XX DR N-PSDB; AAT16516.
XX
XX Chimaeric DNA encoding protein contg. extracellular matrix protein
XX domain - and cellular regulatory factor domain, partic. useful as
XX osteogenic agents, also related vectors, transformed cells and
XX chimaeric proteins.
XX
XX PS Disclosure; Fig 6; 59pp; English.
XX
XX CC A fusion protein (AAR89470) comprises the alpha-helical region of
XX human collagen I(a) linked to the human mature transforming
XX growth factor beta-1 (TGF-beta-1). It can be expressed in
XX Escherichia coli transformants carrying a vector incorporating a
XX chimeric gene (AAT16516) coding for the fusion. The TGF-beta-
XX moiety increases efficacy of the body's normal soft tissue
XX repair response and also induces osteogenesis. The collagen
XX moiety provides an integral substratum or scaffolding for the
XX TGF and cells involved in reconstruction and growth. The fusion
XX protein provides sustained release and delivery of TGF-beta-1
XX to a target tissue.
XX
XX SQ Sequence 1171 AA;
XX
XX Query Match 100.0%; Score 580; DB 17; Length 1171;
XX Best Local Similarity 100.0%; Pred. No. 1.2e-34;
XX Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
XX
QY 1 RGDKGTGEQDRGKIGHRGFSGLOGPPGPGSPGEGQSGAGPAGPPGPGSAGAPGK 60
DB 932 rgdkgtgeqgdrkighrgfsglqgpppgpsgeqgsgagpagrpgppgsagapgk 991
DB
QY 61 DGLNGLPGPIGPPGRGTGDAGVPVGGPPGPPGPPGPP 100
DB 992 dglnlpgpi9ppgrgtgdagvpgppgpppppppp 1031
DB
RESULT 24
ID AAY84538
XX AAY84538 standard; Protein; 1171 AA.
XX
XX AC AAY84538;
XX
XX DT 25-JUL-2000 (first entry)
```

```
XX A chimeric collagen 1 (alphal)/TGF-beta1 protein.
XX Extracellular matrix protein; self aggregation; hydroxylated proline;
XX trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;
XX collagen; fibrinogen; fibronectin; post translational hydroxylation; ss.
XX transforming growth factor-beta1; TGF-beta1; chimera.
XX
XX OS Chimeric - Homo sapiens.
XX OS Chimeric - Unidentified.
XX
XX FH Key Location/Qualifiers
XX FT Misc-difference 858
XX FT /note= "Gly encoded by GCT"
XX
XX PN EP992586-A2.
XX
XX PD 12-APR-2000.
XX
XX PF 07-OCT-1999; 99EP-0119184.
XX
XX PR 09-OCT-1998; 98US-0359768.
XX
XX PA (USSU ) US SURGICAL CORP.
XX
XX PI Gruskin EA, Buechter DD, Zhang G, Connolly K;
XX
XX DR WPI; 2000-259138/23.
XX DR N-PSDB; AAL12498.
XX
XX PT Production of extracellular matrix proteins containing
XX 4-trans-hydroxyproline results in native self aggregating proteins,
XX useful on medical implants -
XX
XX PS Claim 23; Fig 15; 260pp; English.
XX
XX CC The specification describes a method for producing an extracellular
XX matrix protein or its fragment. The extracellular matrix protein is
XX capable of self aggregating in a cell which does not ordinarily
XX hydroxylated prolines. The method comprises optimising a nucleic acid
XX sequence for expression in the cell by substitution of codons preferred
XX by that cell for naturally occurring codons not preferred by the cell;
XX incorporating the nucleic acid sequence into the cell; and contacting
XX the cell with a hypertonic growth medium containing at least one amino
XX acid, selected from the group consisting of trans-4-hydroxyproline and
XX 3-hydroxyproline to allow at least one of the amino acids to be
XX assimilated into the cell and incorporated into the extracellular matrix
XX protein. The method may be used to make host cells assimilate and
XX incorporate trans-4-hydroxyproline into proteins. This is especially
XX useful in the recombinant production of proteins such as collagen,
XX fibrinogen and fibronectin whose ability to self aggregate and produce
XX functional proteins depends on the post translational hydroxylation of
XX proline. The method is also useful in studying the structure and function
XX of polypeptides which do not normally contain trans-4-hydroxyproline.
XX The present sequence represents chimeric collagen 1 (alphal)/transforming
XX growth factor-beta1 (TGF-beta1) protein, which may be produced using the
XX method of the invention.
XX
XX SQ Sequence 1171 AA;
XX
XX Query Match 100.0%; Score 580; DB 21; Length 1171;
XX Best Local Similarity 100.0%; Pred. No. 1.2e-34;
XX Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
XX
QY 1 RGDKGTGEQDRGKIGHRGFSGLOGPPGPGSPGEGQSGAGPAGPPGPGSAGAPGK 60
DB 932 rgdkgtgeqgdrkighrgfsglqgpppgpsgeqgsgagpagrpgppgsagapgk 991
DB
QY 61 DGLNGLPGPIGPPGRGTGDAGVPVGGPPGPPGPPGPP 100
DB 992 dglnlpgpi9ppgrgtgdagvpgppgpppppppp 1031
DB
```

RESULT 25

AAR71701
ID AAR71701 standard; protein; 1341 AA.
XX AC AAR71701;
XX DT 17-OCT-1995 (first entry)
XX DE Collagen alpha 1 (I) chain precursor.
XX KW Collagen; antibody; immunoassay; metabolism; diagnosis; monitoring;
XX KW disorder; osteoporosis; metastatic progression; Paget's disease;
XX KW hyperthyroidism; bone; resorption; rheumatoid arthritis;
XX KW osteoarthritis; vasculitis syndrome.
XX OS Homo sapiens.
XX FH Key Location/Qualifiers
FT Misc-difference 2028 /note= "Unidentified amino acid."
XX WO9508115-A.
XX PN 23-MAR-1995.
XX PD 19-SEP-1994; 94WO-DK00348.
XX PF 17-SEP-1993; 93DK-0001040.
XX PR (OSTE-) OSTEOMETER AS.
XX PA Bonde M, Qvist P;
XX PI WPI; 1995-131456/17.
XX DR Assaying collagen fragments in body fluid by immunoassay - using
PT antibodies raised against synthetic peptide(s) contg. potential
PT crosslinking sites, to diagnose and monitor disorders of collagen
PT metabolism, e.g. osteoporosis.
XX Disclosure (Appendix A); Page 49; 87pp; English.
XX Determination of collagen fragments in body fluids can be achieved
CC by immunoassay using antibodies directed against synthetic peptides
CC derived from collagen which contain sites of potential crosslinking.
CC The method is used to diagnose and monitor treatment of disorders of
CC collagen metabolism (degradation of type I collagen may indicate
CC osteoporosis, metastatic progression, Paget's disease,
CC hyperthyroidism or other conditions involving excessive bone
CC resorption; degradation of type II collagen may indicate rheumatoid
CC arthritis or osteoarthritis, and of type III collagen, vaculitis
CC syndrome). The method can also be used to assess the toxicity of a
CC component of drugs for their effect on collagen metabolism.
XX Sequence 1341 AA;

Query Match 100.0%; Score 580; DB 16; Length 1341;
Best Local Similarity 100.0%; Pred. No. 1.3e-34;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGGQDRGKIHGRFSGLOGPPGPGSGAGSGAGPAGPGGAGAPGK 60
Db 970 rgdkgetgeggdrgikghrgfsglqpppppgsgagsgagpgprgpggsagapgk 1029
QY 61 DGLNGLPGPIGPPGPRGTGDAGVCPGPPGPPGPP 100
Db 1030 dglnglpgpi gpppgprgtgdagvcp gpppgppgpp 1069

RESULT 26

AAY96122
ID AAY96122 standard; Peptide; 1341 AA.
XX AC AAY96122;
XX DT 19-DEC-2000 (first entry)
XX DE Collagen type I alpha-1.
XX KW Collagen type I; osteoporosis; bone resorption; Paget's disease;
XX KW hyperparathyroidism; metastasis; assay; diagnosis.
XX OS Homo sapiens.
XX FH Key Location/Qualifiers
FT Misc-difference 924 /note= "unidentified residue"
FT Misc-difference 927 /note= "unidentified residue"
FT Misc-difference 1127 /note= "unidentified residue"
FT Misc-difference 1268 /note= "unidentified residue"
XX US6110689-A.
XX PN 29-AUG-2000.
XX PD 04-NOV-1997; 97US-0963825.
XX PF 21-JAN-1994; 94US-0187319.
XX PR (OSTE-) OSTEOMETER AS.
XX PA Bonde M, Qvist P;
XX PI WPI; 2000-586349/55.
XX DR Assaying type I collagen fragments for diagnosing osteoporosis in
PT postmenopausal woman, involves contacting body fluid with synthetic
PT collagen peptide and antibody and quantifying by competitive binding
PT assay
XX Disclosure: Column 23-37; 41pp; English.
XX The present sequence is that of human type I collagen alpha-1.
CC The invention is based on the discovery of the presence of
CC particular collagen fragments in body fluids of patients compared
CC with those of healthy subjects. These fragments are generated
CC from collagen degradation and are partly characterised by the
CC presence of potential sites for crosslinking. A method for
CC assaying collagen fragments in a body fluid sample is based on the
CC competitive binding to immunological binding partners of collagen
CC fragments in the sample and of synthetic peptides derived from
CC collagen and containing crosslinkable sites (see AY96105-11). When
CC considering the degradation of type I collagen, the assay can be
CC used as a means of identifying excessive bone resorption, indicating
CC the presence of osteoporosis or the metastatic progress of a
CC malignancy. Other conditions characterized by excessive bone
CC resorption include Paget's disease and hyperparathyroidism.
XX Sequence 1341 AA;

Query Match 100.0%; Score 580; DB 21; Length 1341;
Best Local Similarity 100.0%; Pred. No. 1.3e-34;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGGQDRGKIHGRFSGLOGPPGPGSGAGSGAGPAGPGGAGAPGK 60
Db 969 rgdkgetgeggdrgikghrgfsglqpppppgsgagsgagpgprgpggsagapgk 1028
QY 61 DGLNGLPGPIGPPGPRGTGDAGVCPGPPGPPGPP 100

Db 1029 dglnglpipgppgrgrtdagpvpgppppppppppp 1068

RESULT 27

AAR89471
ID AAR89471 standard; Protein; 1388 AA.

XX AAR89471;

XX 01-OCT-1996 (first entry)

XX Collagen/Decorin fusion protein.

XX Transforming growth factor; TGF-beta-1; collagen IA; osteogenesis;
KW bone formation; tissue repair; fusion protein.

XX Synthetic.

XX Key Location/Qualifiers

FT Domain 1..1057

FT /label= Collagen-IA

FT /note= "collagen IA alpha-helical domain"

FT Peptide 1058..1059

FT /label= Linker_peptide

FT Domain 1060..1388

FT /label= Decorin

FT Misc-difference 887

FT /note= "unidentified amino acid"

FT Misc-difference 890

FT /note= "unidentified amino acid"

XX CA2151547-A.

XX 11-DEC-1995.

XX 12-JUN-1995; 95CA-2151547.

XX 10-JUN-1994; 94US-0259263.

XX (USSU) US SURGICAL CORP.

XX Espino P, Gruskin EA;

XX WPI; 1996-140144/15.

XX N-PSDB; AAT16517.

XX Chimaeric DNA encoding protein contg. extracellular matrix protein
PT domain - and cellular regulatory factor domain, partic. useful as
PT osteogenic agents, also related vectors, transformed cells and
PT chimaeric proteins.

PS Disclosure; Fig 7; 59pp; English.

XX A fusion protein (AAR89471) comprises the alpha-helical region of
CC human collagen I(a) linked to human dermatan sulphate proteoglycan
CC (decorin). It can be expressed in Escherichia coli transformants
CC carrying a vector incorporating a chimeric gene (AAT16517) coding for
CC the fusion. The decorin binds to type I collagen and thus affects
CC Elbril formation. It inhibits the cell attachment-promoting
CC activity of collagen and fibrinogen by binding to such molecules
CC near their cell binding sites. The collagen moiety provides an
CC integral substratum or scaffolding for the decorin. The fusion
CC protein acts to reduce scarring of healing tissue.

XX Sequence 1388 AA;

Query Match 100.0%; Score 580; DB 17; Length 1388;

Best Local Similarity 100.0%; Pred. No. 1.3e-34;

Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 RGDKGTEQGDRIKGRGSGLGQPPGPGPGGPGAGPGRGPGGAGAPGK 60

|||||

Db 932 rgdgetgeqdgrikghrgfsglqgppbpgspgqsgagpagrgppgsagapgk 991

Qy 61 DGLNGLPGPIGPPGPRGRTGDAGPVGPPGPPGPPGPPGPP 100

Db 992 dglnglpipgppgrgrtdagpvpgppppppppppp 1031

RESULT 28

AAY84539

ID AAY84539 standard; Protein; 1388 AA.

XX AAY84539;

XX 25-JUL-2000 (first entry)

XX Amino acid sequence of a chimeric collagen 1 (alpha1)/decorin protein.

XX Extracellular matrix protein; self aggregation; hydroxylated proline;
KW trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;
KW collagen; fibrinogen; fibronectin; post translational hydroxylation;
KW decorin; chimera.

XX Chimeric - Homo sapiens.

XX Chimeric - Unidentified.

XX Key Location/Qualifiers

FT Misc-difference 87

FT /note= "Gly encoded by GCG"

FT Misc-difference 305

FT /note= "Glu encoded by CAA"

FT Misc-difference 363

FT /note= "Gly encoded by CGT"

FT Misc-difference 378

FT /note= "Glu encoded by GGT"

FT Misc-difference 429

FT /note= "Gly encoded by CGA"

FT Misc-difference 444

FT /note= "Gly encoded by CGC"

FT Misc-difference 543

FT /note= "Gly encoded by GCC"

FT Misc-difference 546

FT /note= "Gly encoded by GCT"

FT Misc-difference 606

FT /note= "Gly encoded by GAC"

FT Misc-difference 702

FT /note= "Gly encoded by CGT"

FT Misc-difference 815

FT /note= "Pro encoded by CTT"

FT Misc-difference 858

FT /note= "Gly encoded by GCT"

FT Misc-difference 1066

FT /note= "Gly encoded by GCC"

XX EP992586-A2.

XX 12-APR-2000.

XX 07-OCT-1999; 99EP-0119184.

XX 09-OCT-1998; 98US-0169768.

XX (USSU) US SURGICAL CORP.

XX Gruskin EA, Buechter DD, Zhang G, Connolly K;

XX WPI; 2000-259138/23.

XX N-PSDB; AAA12499.

XX Production of extracellular matrix proteins containing

XX 4-trans-hydroxyproline results in native self aggregating proteins,

XX useful on medical implants -

XX Claim 25; Fig 17A-B; 260pp; English.

XX The specification describes a method for producing an extracellular
CC matrix protein or its fragment. The extracellular matrix protein is
CC capable of self aggregating in a cell which does not ordinarily
CC hydroxylated prolines. The method comprises optimising a nucleic acid
CC sequence for expression in the cell by substitution of codons preferred
CC by that cell for naturally occurring codons not preferred by the cell;
CC incorporating the nucleic acid sequence into the cell; and contacting
CC the cell with a hypertonic growth medium containing at least one amino
CC acid, selected from the group consisting of trans-4-hydroxyproline and
CC 3-hydroxyproline to allow at least one of the amino acids to be
CC assimilated into the cell and incorporated into the extracellular matrix
CC protein. The method may be used to make host cells assimilate and
CC incorporate trans-4-hydroxyproline into proteins. This is especially
CC useful in the recombinant production of proteins such as collagen,
CC fibrinogen and fibronectin whose ability to self aggregate and produce
CC functional proteins depends on the post translational hydroxylation of
CC proline. The method is also useful in studying the structure and function
CC of polypeptides which do not normally contain trans-4-hydroxyproline.
CC The present sequence represents a chimeric collagen I (alpha1)/decorin
CC protein, which may be produced using the method of the invention.
XX
SQ Sequence 1388 AA;

Query Match 100.0%; Score 580; DB 21; Length 1388;
Best Local Similarity 100.0%; Pred. No. 1.3e-34;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGGQDRGKIKHGRFSGLOGPPGPGSGFGQPSGAGPAGPRGPGSAGAPGK 60
DB 932 rgdkgtgeggdrgikghrgfsglqpppppgpsgsgagpgprgpggsagapgk 991
QY 61 DGLNGLPGPIGPPGRGTGDAGVPGPPGPPGPPGPP 100
DB 992 dginglpgpigrgrtgdagvpgpppgpppgpppp 1031

RESULT 29
AAE02535
ID AAE02535 standard; Protein; 1449 AA.
AC AAE02535;
XX
DT 10-AUG-2001 (first entry)
XX
DE Porcine alpha1(I) collagen.
XX
KW Porcine; alpha1(I) collagen; gelatin; cytostatic; viral infection;
KW pharmaceutical; food industry; cosmetic; autoimmune disorder; vaccine;
KW medical; arterial sealant; bone graft; dermal implant; haemostat; cancer;
KW rheumatoid arthritis; beverage; photographic application.
XX
OS Sus scrofa.
XX
FH Key Location/Qualifiers
FT Misc-difference 829..830
FT /note= "Encoded by ggcaaaccttggtgatctggtgctaaaggcgtg
FT ctggtcccccgcctgctgga"
XX
PN WO200134647-A2.
XX
PD 17-MAY-2001.
XX
PF 10-NOV-2000; 2000WO-US30792.
XX
PR 12-NOV-1999; 99US-0439058.
PR 10-NOV-2000; 2000US-0439058.
XX
PA (FIBR-) FIBROGEN INC.
XX
PI Bell MP, Neff TB, Polarek JW, Seeley TW;
XX

DR WPI; 2001-335911/35.
XX N-PSDB; AAD06576.
PT Novel isolated and purified bovine or porcine collagens and gelatins
PT useful in medical, pharmaceutical, food and cosmetic industries, as
PT vaccine, and for treating autoimmune disorders, infections and cancer
XX
XX Example 3; Fig 8; 168pp; English.
XX The present sequence is porcine alpha1(I) collagen. The present
CC invention relates to recombinant synthesis of collagens and gelatins
CC derived from animals. Collagen is useful in medical, pharmaceutical,
CC food and cosmetic industries. Collagen is an important component of
CC arterial sealants, bone grafts, drug delivery system, dermal implants,
CC haemostats, and incontinence implants, and for treating autoimmune
CC disorders such as rheumatoid arthritis. Collagen is useful in food
CC products such as sausage casings, and in cosmetics or facial and skin
CC products such as moisturisers. Recombinant gelatin is useful in vaccine
CC formulations for treating viral infections, autoimmune diseases and
CC cancer. Gelatin is useful in the manufacture or as a component of
CC various pharmaceutical and medical devices and products, in food and
CC beverage industries, in hair care and skin care products, as a glue or
CC adhesive in various manufacturing processes, as a light-sensitive coating
CC in various electronic devices, as photoresist base in photolithographic
CC processes, in printing and photographic applications, in laboratory
CC application, and as a component in various gels used for biochemical and
CC electrophoretic analysis, including enzymographic gels.
XX
SQ Sequence 1449 AA;
Query Match 100.0%; Score 580; DB 22; Length 1449;
Best Local Similarity 100.0%; Pred. No. 1.4e-34;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 RGDKGTGGQDRGKIKHGRFSGLOGPPGPGSGFGQPSGAGPAGPRGPGSAGAPGK 60
DB 1078 rgdkgtgeggdrgikghrgfsglqpppppgpsgsgagpgprgpggsagapgk 1137
QY 61 DGLNGLPGPIGPPGRGTGDAGVPGPPGPPGPPGPP 100
DB 1138 dginglpgpigrgrtgdagvpgpppgpppgpppp 1177

RESULT 30
AAW68485
ID AAW68485 standard; Protein; 1464 AA.
XX
AC AAW68485;
XX
DT 08-DEC-1998 (first entry)
XX
DE Human recombinant collagen protein.
XX
KW Primer; PCR; amplification; human; collagen; mammal; plant; prosthesis;
KW cardiac valve; ligament; tendon; skin; gingival implant; perfumes;
KW nerve regeneration; antibiotic; growth factor; cancer; inflammatory;
KW gelatin; glue; food.
XX
OS Synthetic.
OS Homo sapiens.
XX
FH Key Location/Qualifiers
FT Peptide 1..22
FT /note= "signal peptide"
FT Protein 23..999
FT /note= "mature protein"
FT Cleavage-site 161
FT /note= "cleavage site for aminopeptidase"
FT Cleavage-site 1218
FT /note= "cleavage site for carboxypeptidase"
XX

PN WO9827202-A1.
XX 25-JUN-1998.
XX 17-DEC-1997; 97WO-FR02331.
XX 17-DEC-1996; 96FR-0016224.
XX (BIOC-) BIOCEM SA.
XX Bournat P, Comte J, Exposito JV, Garrone R, Gruber V;
PI Merot B, Ruggiero F;
XX WPI; 1998-362771/31.
DR N-PSDB; AAV60814.
XX
PT New recombinant nucleic acid for expressing collagen or derivatives
in plants - useful as, e.g. bio-materials and in therapeutic,
PT cosmetic and odontological compositions
XX
PS Disclosure; Fig 7; 138pp; French.
XX
CC The invention relates to the production of mammalian collagen in plants.
CC 2 clones: alpha3 and alpha22, spanning the human collagen type I gene
CC were isolated from a MG-63 osteosarcoma library. Clone alpha3 contained
CC 83 bp of the 5' untranslated region and the first 1920 bp of coding
CC sequence, whereas clone alpha22 contained sequence encoding amino acids
CC 171-1454 of the protein and around 500 bp of the 3' untranslated region.
CC The 2 clones were used to generate a number of fragments which were used
CC to construct composite sequences encoding variant collagen molecules.
CC The fragments are: (A) containing nucleotides (nt) -4 to 479;
CC (B) containing TAA upstream of the sequence encoding the PRS
CC (pathogenesis-related protein S) signal peptide and bases 66-77 from the
CC sequence encoding the N-terminus of the pro-collagen amino propeptide
CC domain; (C) the whole of the amino propeptide domain (nt 72-479); (D) all
CC of the amino-telopeptide domain (nt 474-534); and the N-terminus of the
CC helical region (nt 535-1920); (E) the DRAIII-BamHI fragment
CC (1709-2808) of alpha22, encoding aa 567-936 of the central helical
CC domain; (F) the BamHI-EcoRI (2803-4362) region of alpha22, encoding
CC aa 936-1192 in the central helical domain and aa 1193-1454 in the
CC C-propeptide domain; (G) the C-terminus of the C-propeptide domain
CC (aa 1346-1464) plus stop codons, and (H) as G but encoding aa 1343-1401
CC and also including the KDEI motif for retention in the ER. This sequence
CC represents a recombinant human collagen. The encoding gene was
CC constructed from fragments (A), (D), (E), (F) and (G). The recombinant
CC gene is used for expression of mammalian collagen in plant cells. The
CC transformed plants, their extracts and parts are useful as biomaterials
CC (haemostatic compresses, sponges or bandages) and in pharmaceutical,
CC medical, odontological, cosmetic and biotechnological compositions (e.g.
CC as prostheses for cardiac valves, ligaments or tendons; skin substitutes;
CC gingival implants; microcapsules for perfumes; guide tubes for nerve
CC regeneration; slow release products for antibiotics, growth factors,
CC anticancer agents or anti-inflammatories; surgical thread and components
CC of ointments). They are suitable for treating any disorder related to
CC collagen dysfunction and gelatin, produced from collagen, is used to
CC produce gylues, surgical prostheses and foods.
XX
SQ Sequence 1464 AA;

Query Match 100.0%; Score 580; DB 19; Length 1464;
Best Local Similarity 100.0%; Pred. No. 1.4e-34;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 RGDGKGTGCGDRIKGRGSGLGQPPGSGCGECPGASGAPGPPGSGAGPCK 60
|||||
Db 1093 rgdkgetegdgdkgrgsglgqppgpgsgcgsgagpgrppgsagapgk 1152

Qy 61 DGLNGLPGIPGPPGPRGTGAGVGPVGPVGPVGPVGPVGPVGPV 100
|||||

Db 1153 dglnglpipgpppgrgtgagvgppvgppvgppvgppvgppvgpp 1192

RESULT 31
AAU14136
ID AAU14136 standard; Protein; 1464 AA.
XX AC AAU14136;
XX 24-OCT-2001 (first entry)
XX Human novel protein #7.
XX
XX Human; novel protein; Antianaemic; osteopathic; antiinflammatory;
KW immunomodulatory; cytostatic; neuroprotective; vulnerary; nootropic;
KW anticonvulsant; antiarthritic; cerebroprotective; vulnerary; antifungal; antiviral;
KW antibacterial; antiallergic; dermatological; haemostatic; antiasthmatic;
KW thrombolytic; immunogen; antibody; gene therapy; neurological disorder;
KW Parkinson's disease; inflammatory disorder; cancer; asthma; osteoporosis;
XX tissue regeneration; immune disorder.
OS Homo sapiens.
XX WO200155437-A2.
XX 02-AUG-2001.
XX 25-JAN-2001; 2001WO-US02623.
XX 25-JAN-2000; 2000US-0491404.
XX (HYSE-) HYSEQ INC.
XX Tang YT, Liu C, Drmanac RT;
PI WPI; 2001-451939/48.
DR N-PSDB; AAS22441.
XX Isolated polypeptides useful for treating anti-inflammatory diseases,
PT nervous system disorders, and for regenerating bone and cartilage -
XX Example 4; Page 525-527; 894pp; English.
XX The invention relates to polynucleotides encoding novel human
CC proteins or their active domains. The polypeptides, polynucleotides and
CC antibodies raised against the polypeptides are used in a method of
CC treatment of a mammal and prevention of disorders caused by the aberrant
CC protein expression or activity. The polypeptides can be used as
CC molecular weight markers, food supplements, and in antibody production.
CC The polypeptides are used to identify compounds which bind to the
CC polypeptides. Polynucleotides of the invention are used as probes and
CC primers, for sequencing, for chromosome or gene mapping, in the
CC production of recombinant proteins, and in generating anti-sense DNA or
CC RNA and in gene therapy. Polypeptides of the invention can be used to
CC target drugs to a tumour, in assays to determine biological activity, to
CC raise antibodies/elicit an immune response, to determine quantitative
CC protein levels, as tissue markers, and to isolate receptors or ligands.
CC Polypeptides of the invention may also be useful in treating platelet
CC disorders, stem cell disorders, regenerating bone, cartilage, tendon,
CC ligament and/or nerve tissue, wound healing, treating burns, promoting
CC the proliferation, differentiation and survival of stem cells, as a
CC contraceptive, treating osteoporosis and osteoarthritis, anaemia, as a
CC Alzheimer's, Parkinson's and Huntington's diseases, amyotrophic lateral
CC sclerosis, stroke, immune deficiencies resulting from bacterial, viral or
CC fungal infection or from autoimmunity, cancer, allergy, asthma,
CC graft-versus-host disease, eczema, haemophilia, thrombosis,
CC anti-inflammatory diseases, nervous system disorders, and infection.
XX The present sequence represents a protein of the invention.

Sequence 1464 AA;

Query Match 100.0%; Score 580; DB 22; Length 1464;
Best Local Similarity 100.0%; Pred. No. 1.4e-34;
Matches 100; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

GenCore version 4.5
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OM protein - protein search, using sw model

Run on: January 29, 2002, 12:14:46 ; Search time 38.4 Seconds
(without alignments)
113.811 Million cell updates/sec

Title: US-09-710-239-18

Perfect score: 333

Sequence: 1 EAGLPCAKGLTSPGSPD.....PPCARGQACVGMFGPGKGA 59

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 522463 seqs, 74073290 residues

Total number of hits satisfying chosen parameters: 522463

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

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- 1: /SID58/gcgdata/geneseq/geneseq/AA1980.DAT:*
- 2: /SID58/gcgdata/geneseq/geneseq/AA1981.DAT:*
- 3: /SID58/gcgdata/geneseq/geneseq/AA1982.DAT:*
- 4: /SID58/gcgdata/geneseq/geneseq/AA1983.DAT:*
- 5: /SID58/gcgdata/geneseq/geneseq/AA1984.DAT:*
- 6: /SID58/gcgdata/geneseq/geneseq/AA1985.DAT:*
- 7: /SID58/gcgdata/geneseq/geneseq/AA1986.DAT:*
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- 10: /SID58/gcgdata/geneseq/geneseq/AA1989.DAT:*
- 11: /SID58/gcgdata/geneseq/geneseq/AA1990.DAT:*
- 12: /SID58/gcgdata/geneseq/geneseq/AA1991.DAT:*
- 13: /SID58/gcgdata/geneseq/geneseq/AA1992.DAT:*
- 14: /SID58/gcgdata/geneseq/geneseq/AA1993.DAT:*
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- 19: /SID58/gcgdata/geneseq/geneseq/AA1998.DAT:*
- 20: /SID58/gcgdata/geneseq/geneseq/AA1999.DAT:*
- 21: /SID58/gcgdata/geneseq/geneseq/AA2000.DAT:*
- 22: /SID58/gcgdata/geneseq/geneseq/AA2001.DAT:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	333	100.0	59	22	AAE02704 Human alpha (I) t
2	333	100.0	59	22	AAE02704 Amino acid sequenc
3	333	100.0	101	22	AAE02705 Human alpha (I) t
4	333	100.0	101	22	AAE02705 Amino acid sequenc
5	333	100.0	185	22	AAE02706 Human alpha (I) t
6	333	100.0	185	22	AAE02706 Amino acid sequenc
7	333	100.0	251	22	AAE02707 Human alpha (I) t
8	333	100.0	251	22	AAE02707 Amino acid sequenc
9	333	100.0	500	22	AAE02708 Human alpha (I) t
10	333	100.0	500	22	AAE02708 Amino acid sequenc
11	333	100.0	501	22	AAE02703 Human alpha (I) t

12	333	100.0	501	22	AAE02704 Human alpha (I) t
13	333	100.0	662	22	AAE02718 Amino acid sequenc
14	333	100.0	662	22	AAE02718 Amino acid sequenc
15	333	100.0	1057	21	AAE02718 Amino acid sequenc
16	333	100.0	1057	21	AAE02718 Amino acid sequenc
17	333	100.0	1058	21	AAE02718 Amino acid sequenc
18	333	100.0	1107	17	AAE02718 Amino acid sequenc
19	333	100.0	1107	17	AAE02718 Amino acid sequenc
20	333	100.0	1169	17	AAE02718 Amino acid sequenc
21	333	100.0	1169	17	AAE02718 Amino acid sequenc
22	333	100.0	1171	17	AAE02718 Amino acid sequenc
23	333	100.0	1171	17	AAE02718 Amino acid sequenc
24	333	100.0	1341	16	AAE02718 Amino acid sequenc
25	333	100.0	1341	16	AAE02718 Amino acid sequenc
26	333	100.0	1388	17	AAE02718 Amino acid sequenc
27	333	100.0	1411	21	AAE02718 Amino acid sequenc
28	333	100.0	1449	22	AAE02718 Amino acid sequenc
29	333	100.0	1463	22	AAE02718 Amino acid sequenc
30	333	100.0	1464	19	AAE02718 Amino acid sequenc
31	333	100.0	1464	22	AAE02718 Amino acid sequenc
32	333	100.0	1464	22	AAE02718 Amino acid sequenc
33	325	97.6	1388	21	AAE02718 Amino acid sequenc
34	321	96.4	595	20	AAE02718 Amino acid sequenc
35	321	96.4	822	20	AAE02718 Amino acid sequenc
36	256	76.9	1418	15	AAE02718 Amino acid sequenc
37	256	76.9	1418	16	AAE02718 Amino acid sequenc
38	256	76.9	1418	21	AAE02718 Amino acid sequenc
39	256	76.9	1418	22	AAE02718 Amino acid sequenc
40	256	76.9	1442	16	AAE02718 Amino acid sequenc
41	256	76.9	1487	19	AAE02718 Amino acid sequenc
42	234	70.3	1078	16	AAE02718 Amino acid sequenc
43	234	70.3	1078	21	AAE02718 Amino acid sequenc
44	234	70.3	1196	13	AAE02718 Amino acid sequenc
45	234	70.3	1466	22	AAE02718 Amino acid sequenc

ALIGNMENTS

RESULT	1
AAE02704	AAE02704 standard; Protein; 59 AA.
ID	AAE02704; AC
AC	AAE02704; DT
DT	06-AUG-2001 (first entry)
DE	Human alpha (I) type I collagen helical domain (residues 531-589).
XX	Human; recombinant gelatin; binding agent; stabilising agent; emulsifier; encapsulant; film-forming agent; moisturising agent; thickening agent; gelling agent; colloidal agent; adhesive agent; gel capsule; photography; plasma expander; colloidal volume replacement material; graft coating; medical sponge; medical plug; micro-carrier; edible composition; protein supplement; fat substitute; nutritional supplement; cell culture; edible coating; cosmetic; vaccine; therapy; arthritis; attheros; cartilage degeneration; joint flexibility; food industry; beverage; alpha (I) type I collagen.
XX	Homo sapiens.
OS	WO200134646-A2.
PN	17-MAY-2001.
PD	10-NOV-2000; 2000WO-US30791.
PF	12-NOV-1999; 99US-0165114.
PR	15-MAY-2000; 2000US-0204437.
PA	(FIBR-) FIBROGEN INC.
PI	Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;

XX WPI; 2001-329072/34.
XX
XX Gelatin useful for pharmaceuticals, cosmetics and edible foods, is
PT prepared recombinantly -
XX
XX Claim 21; Page 123; 137pp; English.
XX
XX The patent discloses recombinant human gelatin which is useful
CC in various compositions including binding agents, encapsulants,
CC stabilising agents, film-forming agents, moisturising agents,
CC emulsifiers, thickening agents, gelling agents, colloidal agents,
CC adhesive agents, pharmaceutical compositions, hard gel capsules,
CC soft gel capsules, plasma expander, colloidal volume replacement
CC materials, graft coatings, medical sponges, medical plugs,
CC pharmaceutical stabilisers, micro-carriers, edible compositions,
CC protein supplements, fat substitutes, nutritional supplements,
CC edible coatings, photographic compositions, cosmetic compositions,
CC industrial composition, cell culture compositions and compositions
CC for use in the laboratory. Pharmaceutical compositions comprising
CC recombinant gelatin are used as vaccines. They are also used to
CC treat various joint conditions such as arthritis, athrosis and
CC other conditions related to the degeneration of cartilage and joint
CC flexibility. Recombinant gelatin is also used in food and beverage
CC industries. The present sequence is human alpha (I) type I collagen
CC helical domain (residues 531-589). This sequence is a recombinant
CC gelatin.
XX
XX Sequence 59 AA;
SQ

Query Match 100.0%; Score 333; DB 22; Length 59;
Best Local Similarity 100.0%; Pred. No. 5.6e-24;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 EAGLPKAKGLTSGSPGPDGKTPGPPGAGQDGRPGPPGARGQAGVMGFPKGA 59
DB 1 eaglpkagltsgspgpdgktpgppgagqdgrrpgppgargqagvmgfpkgaa 59

RESULT 2
AAB68058
ID AAB68058 standard; Protein; 59 AA.
XX
XX AAB68058;
XX
XX 09-JUL-2001 (first entry)
XX
XX Amino acid sequence of a recombinant human gelatin.
XX
XX Human; gelatin; vaccine; anaphylactic reaction.
XX
XX Homo sapiens.
XX
XX WO200134801-A2.
XX
XX 17-MAY-2001.
XX
XX 10-NOV-2000; 2000WO-US30843.
XX
XX 12-NOV-1999; 99US-0165114.
XX
XX 15-MAY-2000; 2000US-0204437.
XX
XX (FIBR-) FIBROGEN INC.
XX
XX Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX WPI; 2001-308784/32.
XX

XX Vaccine formulations (I) comprising recombinant human gelatin, useful
PT for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies
PT and cholera, the gelatin is non-immunogenic and confers stability at
PT ambient temperatures -

XX Claim 11; Page 116; 130pp; English.
XX
XX The present sequence represents a human recombinant gelatin polypeptide.
CC The recombinant gelatin polypeptide is used to produce vaccine
CC formulations of the invention. The recombinant human gelatin is
CC non-immunogenic (therefore reducing anaphylactic reactions) and confers
CC stability at ambient temperatures. The vaccine formulation comprises a
CC vaccine formulated for the prevention of a disease selected from vaccinia
CC virus (small pox), polio virus (Salk and Sabin), mumps, measles, rubella,
CC diphtheria, tetanus, Varicella-Zoster (chicken pox/shingles), pertussis
CC (whooping cough), Bacille Calmette-Guérin (BCG, tuberculosis),
CC haemophilus influenzae meningitis, rabies, cholera, Japanese
CC encephalitis virus, salmonella typhi, shigella, hepatitis A and B,
CC adenovirus, yellow fever, foot and mouth disease, herpes simplex virus,
CC respiratory syncytial virus, rotavirus, Dengue, West Nile virus, turkey
CC herpes virus (Marek's disease), influenza and/or anthrax.
XX
XX Sequence 59 AA;
SQ

Query Match 100.0%; Score 333; DB 22; Length 59;
Best Local Similarity 100.0%; Pred. No. 5.6e-24;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 EAGLPKAKGLTSGSPGPDGKTPGPPGAGQDGRPGPPGARGQAGVMGFPKGA 59
DB 1 eaglpkagltsgspgpdgktpgppgagqdgrrpgppgargqagvmgfpkgaa 59

RESULT 3
AAE02705
ID AAE02705 standard; Protein; 101 AA.
XX
XX AAE02705;
XX
XX 06-AUG-2001 (first entry)
XX
XX Human alpha (I) type I collagen helical domain (residues 531-631).
XX
XX Human; recombinant gelatin; binding agent; stabilising agent; emulsifier;
XX encapsulant; film-forming agent; moisturising agent; thickening agent;
XX gelling agent; colloidal agent; adhesive agent; gel capsule; photography;
XX plasma expander; colloidal volume replacement material; graft coating;
XX medical sponge; medical plug; micro-carrier; edible composition;
XX protein supplement; fat substitute; nutritional supplement; cell culture;
XX edible coating; cosmetic; vaccine; therapy; arthritis; athrosis;
XX cartilage degeneration; joint flexibility; food industry; beverage;
XX alpha (I) type I collagen.
XX
XX Homo sapiens.
XX
XX WO200134646-A2.
XX
XX 17-MAY-2001.
XX
XX 10-NOV-2000; 2000WO-US30791.
XX
XX 12-NOV-1999; 99US-0165114.
XX
XX 15-MAY-2000; 2000US-0204437.
XX
XX (FIBR-) FIBROGEN INC.
XX
XX Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX WPI; 2001-329072/34.
XX

XX Gelatin useful for pharmaceuticals, cosmetics and edible foods, is
PT prepared recombinantly -
XX
XX Claim 21; Page 123-124; 137pp; English.
XX
XX The patent discloses recombinant human gelatin which is useful

CC in various compositions including binding agents, encapsulants,
 CC stabilising agents, film-forming agents, moisturising agents,
 CC emulsifiers, thickening agents, gelling agents, colloidal agents,
 CC adhesive agents, pharmaceutical compositions, hard gel capsules,
 CC soft gel capsules, plasma expander, colloidal volume replacement
 CC materials, graft coatings, medical sponges, medical plugs,
 CC pharmaceutical stabilisers, micro-carriers, edible compositions,
 CC protein supplements, fat substitutes, nutritional supplements,
 CC edible coatings, photographic compositions, cosmetic compositions,
 CC industrial composition, cell culture compositions and compositions
 CC for use in the laboratory. Pharmaceutical compositions comprising
 CC recombinant gelatin are used as vaccines. They are also used to
 CC treat various joint conditions such as arthritis, atrosis and
 CC other conditions related to the degeneration of cartilage and joint
 CC flexibility. Recombinant gelatin is also used in food and beverage
 CC industries. The present sequence is human alpha (I) type I collagen
 CC helical domain (residues 531-631). This sequence is a recombinant
 CC gelatin.

XX
 SQ Sequence 101 AA;

Query Match 100.0%; Score 333; DB 22; Length 101;
 Best Local Similarity 100.0%; Pred. No. 9.3e-24;
 Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 EAGLPAGKGLTSGSPGDKTGPAGDGRPPGPPGARGAGVGMFGPKCAA 59
 |||||
 DB 1 eaglpagkgltspspgdgktgppagagdgrrpppggargagvngfpgpkga 59

RESULT 4
 AAB68059
 ID AAB68059 standard; Protein: 101 AA.
 XX
 AC AAB68059;
 XX
 DT 09-JUL-2001 (first entry)
 XX
 DE Amino acid sequence of a recombinant human gelatin.
 XX
 KW Human; gelatin; vaccine; anaphylactic reaction.
 XX
 OS Homo sapiens.
 XX
 PN WO200134801-A2.
 XX
 PD 17-MAY-2001.
 XX
 PF 10-NOV-2000; 2000WO-US30843.
 XX
 PR 12-NOV-1999; 99US-0165114.
 PR
 PA 15-MAY-2000; 2000US-0204437.
 XX
 (FIBR-) FIBROGEN INC.
 PA
 PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
 XX
 XX WPI; 2001-308784/32.
 DR
 XX
 PT Vaccine formulations (I) comprising recombinant human gelatin, useful
 PT for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies
 PT and cholera, the gelatin is non-immunogenic and confers stability at
 PT ambient temperatures -
 XX
 XX Claim 11; Page 116-117; 130pp; English.
 PS
 CC The present sequence represents a human recombinant gelatin polypeptide.
 CC The recombinant gelatin polypeptide is used to produce vaccine
 CC formulations of the invention. The recombinant human gelatin is
 CC non-immunogenic (therefore reducing anaphylactic reactions) and confers
 CC stability at ambient temperatures. The vaccine formulation comprises a
 CC vaccine formulated for the prevention of a disease selected from vaccinia

CC virus (small pox), polio virus (Salk and Sabin), mumps, measles, rubella,
 CC diphtheria, tetanus, Varicella-Zoster (chicken pox/shingles), pertussis
 CC (whooping cough), Bacille Calmette-Guérin (BCG, tuberculosis),
 CC haemophilus influenzae meningitis, rabies, cholera, Japanese
 CC encephalitis virus, salmonella typhi, shigella, hepatitis A and B,
 CC adenovirus, yellow fever, foot and mouth disease, herpes simplex virus,
 CC respiratory syncytial virus, rotavirus, Dengue, West Nile virus, turkey
 CC herpes virus (Marek's disease), influenza and/or anthrax.

XX
 SQ Sequence 101 AA;

Query Match 100.0%; Score 333; DB 22; Length 101;
 Best Local Similarity 100.0%; Pred. No. 9.3e-24;
 Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 EAGLPAGKGLTSGSPGDKTGPAGDGRPPGPPGARGAGVGMFGPKCAA 59
 |||||
 DB 1 eaglpagkgltspspgdgktgppagagdgrrpppggargagvngfpgpkga 59

RESULT 5
 AAE02706
 ID AAE02706 standard; Protein: 185 AA.
 XX
 AC AAE02706;
 XX
 DT 06-AUG-2001 (first entry)
 XX
 DE Human alpha (I) type I collagen helical domain (residues 531-715).
 XX
 KW Human; recombinant gelatin; binding agent; stabilising agent; emulsifier;
 KW encapsulant; film-forming agent; moisturising agent; thickening agent;
 KW gelling agent; colloidal agent; adhesive agent; gel capsule; photography;
 KW plasma expander; colloidal volume replacement material; graft coating;
 KW medical sponge; medical plug; micro-carrier; edible composition;
 KW protein supplement; fat substitute; nutritional supplement; cell culture;
 KW edible coating; cosmetic; vaccine; therapy; arthritis; atrosis;
 KW cartilage degeneration; joint flexibility; food industry; beverage;
 KW alpha (I) type I collagen.
 XX
 OS Homo sapiens.
 XX
 PN WO200134646-A2.
 XX
 PD 17-MAY-2001.
 XX
 PF 10-NOV-2000; 2000WO-US30791.
 XX
 PR 12-NOV-1999; 99US-0165114.
 PR
 PA 15-MAY-2000; 2000US-0204437.
 XX
 (FIBR-) FIBROGEN INC.
 PA
 PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
 XX
 XX WPI; 2001-329072/34.
 DR
 XX
 PT Gelatin useful for pharmaceuticals, cosmetics and edible foods, is
 PT prepared recombinantly -
 XX
 XX Claim 21; Page 124; 137pp; English.
 PS
 CC The patent discloses recombinant human gelatin which is useful
 CC in various compositions including binding agents, encapsulants,
 CC stabilising agents, film-forming agents, moisturising agents,
 CC emulsifiers, thickening agents, gelling agents, colloidal agents,
 CC adhesive agents, pharmaceutical compositions, hard gel capsules,
 CC soft gel capsules, plasma expander, colloidal volume replacement
 CC materials, graft coatings, medical sponges, medical plugs,
 CC pharmaceutical stabilisers, micro-carriers, edible compositions,
 CC protein supplements, fat substitutes, nutritional supplements,
 CC edible coatings, photographic compositions, cosmetic compositions,

CC industrial composition, cell culture compositions and compositions
CC for use in the laboratory. Pharmaceutical compositions comprising
CC recombinant gelatin are used as vaccines. They are also used to
CC treat various joint conditions such as arthritis, athrosis and
CC other conditions related to the degeneration of cartilage and joint
CC flexibility. Recombinant gelatin is also used in food and beverage
CC industries. The present sequence is human alpha (I) type I collagen
CC helical domain (residues 531-715). This sequence is a recombinant
CC gelatin.
XX
SQ Sequence 185 AA;

Query Match 100.0%; Score 333; DB 22; Length 185;
Best Local Similarity 100.0%; Pred. No. 1.6e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 EAGLPGAKGLTSGSPGDPGKGTGPPGAGQDGRPGPGARGQAGVMGFPgpkGAA 59
|||||
Db 1 eaglpgakltsgspgdpdgtgppgagdgrrppgppgargagvmgfpgpkga 59
|||||

RESULT 6
AAB68060
ID AAB68060 standard; Protein: 185 AA.
XX
AC AAB68060;
XX
DT 09-JUL-2001 (first entry)
XX
DE Amino acid sequence of a recombinant human gelatin.
XX
KW Human; gelatin; vaccine; anaphylactic reaction.
XX
OS Homo sapiens.
XX
PN WO200134801-A2.
XX
PD 17-MAY-2001.
XX
PF 10-NOV-2000; 2000WO-US30843.
XX
PR 12-NOV-1999; 99US-0165114.
PR 15-MAY-2000; 2000US-0204437.
XX
PA (FIBR-) FIBROGEN INC.
XX
PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX
DR WPI; 2001-308784/32.
XX
PT Vaccine formulations (I) comprising recombinant human gelatin, useful
PT for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies
PT and cholera, the gelatin is non-immunogenic and confers stability at
PT ambient temperatures -
XX
PS Claim 11; Page 117; 130pp; English.
XX
CC The present sequence represents a human recombinant gelatin polypeptide.
CC The recombinant gelatin polypeptide is used to produce vaccine
CC formulations of the invention. The recombinant human gelatin is
CC non-immunogenic (therefore reducing anaphylactic reactions) and confers
CC stability at ambient temperatures. The vaccine formulation comprises a
CC vaccine formulated for the prevention of a disease selected from vaccinia
CC virus (small pox), polio virus (Salk and Sabin), mumps, measles, rubella,
CC diphtheria, tetanus, Varicella-Zoster (chicken pox/shingles), pertussis
CC (whooping cough), Bacille Calmette-Guérin (BCG, tuberculosis),
CC haemophilus influenzae meningitis, rabies, cholera, Japanese
CC encephalitis virus, salmonella typhi, shigella, hepatitis A and B,
CC adenovirus, yellow fever, foot and mouth disease, herpes simplex virus,
CC respiratory syncytial virus, rotavirus, Dengue, West Nile virus, turkey
CC herpes virus (Marek's disease), influenza and/or anthrax.
XX

SQ Sequence 185 AA;

Query Match 100.0%; Score 333; DB 22; Length 185;
Best Local Similarity 100.0%; Pred. No. 1.6e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 EAGLPGAKGLTSGSPGDPGKGTGPPGAGQDGRPGPGARGQAGVMGFPgpkGAA 59
|||||
Db 1 eaglpgakltsgspgdpdgtgppgagdgrrppgppgargagvmgfpgpkga 59
|||||

RESULT 7
AAE02707
ID AAE02707 standard; Protein: 251 AA.
XX
AC AAE02707;
XX
DT 06-AUG-2001 (first entry)
XX
DE Human alpha (I) type I collagen helical domain (residues 531-781).
XX
KW Human; recombinant gelatin; binding agent; stabilising agent; emulsifier;
KW encapsulant; film-forming agent; moisturing agent; thickening agent;
KW gelling agent; colloidal agent; adhesive agent; gel capsule; photography;
KW plasma expander; colloidal volume replacement material; graft coating;
KW medical sponge; medical plug; micro-carrier; edible composition;
KW protein supplement; fat substitute; nutritional supplement; cell culture;
KW edible coating; cosmetic; vaccine; therapy; arthritis; attheros;
KW cartilage degeneration; joint flexibility; food industry; beverage;
KW alpha (I) type I collagen.
XX
OS Homo sapiens.
XX
PN WO200134646-A2.
XX
PD 17-MAY-2001.
XX
PF 10-NOV-2000; 2000WO-US30791.
XX
PR 12-NOV-1999; 99US-0165114.
PR 15-MAY-2000; 2000US-0204437.
XX
PA (FIBR-) FIBROGEN INC.
XX
PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX
DR WPI; 2001-329072/34.
XX
PT Gelatin useful for pharmaceuticals, cosmetics and edible foods, is
PT prepared recombinantly -
XX
PS Claim 21; Page 125; 137pp; English.
XX
CC The patent discloses recombinant human gelatin which is useful
CC in various compositions including binding agents, encapsulants,
CC stabilising agents, film-forming agents, moisturing agents,
CC emulsifiers, thickening agents, gelling agents, colloidal agents,
CC adhesive agents, pharmaceutical compositions, hard gel capsules,
CC soft gel capsules, plasma expander, colloidal volume replacement
CC materials, graft coatings, medical sponges, medical plugs,
CC pharmaceutical stabilisers, micro-carriers, edible compositions,
CC protein supplements, fat substitutes, nutritional supplements,
CC edible coatings, photographic compositions, cosmetic compositions,
CC industrial composition, cell culture compositions and compositions
CC for use in the laboratory. Pharmaceutical compositions comprising
CC recombinant gelatin are used as vaccines. They are also used to
CC treat various joint conditions such as arthritis, athrosis and
CC other conditions related to the degeneration of cartilage and joint
CC flexibility. Recombinant gelatin is also used in food and beverage
CC industries. The present sequence is human alpha (I) type I collagen
CC helical domain (residues 531-781). This sequence is a recombinant
CC gelatin.
CC

XX SQ Sequence 251 AA;

Query Match 100.0%; Score 333; DB 22; Length 251;
Best Local Similarity 100.0%; Pred. No. 2.2e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 EAGLPAGKGLTGSPPGDKTGPAGQDGRPPGPPGARGQAGVMGFPKGA 59
|||||
Db 1 eaglpagkltgspgspdgktgppagdqgrppgppgargqagvmfpgpkga 59

RESULT 8

AAB68061
ID AAB68061 standard; Protein: 251 AA.
AC AAB68061;
DT 09-JUL-2001 (first entry)
DE Amino acid sequence of a recombinant human gelatin.
KW Human; gelatin; vaccine; anaphylactic reaction.
OS Homo sapiens.
PN WO200134801-A2.
PD 17-MAY-2001.

XX 10-NOV-2000; 2000WO-US30843.
XX 12-NOV-1999; 99US-0165114.
XX 15-MAY-2000; 2000US-0204437.
XX (FIBR-) FIBROGEN INC.
XX Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX WPI; 2001-308784/32.

XX Vaccine formulations (I) comprising recombinant human gelatin, useful
XX for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies
XX and cholera, the gelatin is non-immunogenic and confers stability at
XX ambient temperatures -
XX Claim 11; Page 118; 130pp; English.

XX The present sequence represents a human recombinant gelatin polypeptide.
XX The recombinant gelatin polypeptide is used to produce vaccine
XX formulations of the invention. The recombinant human gelatin is
XX non-immunogenic (therefore reducing anaphylactic reactions) and confers
XX stability at ambient temperatures. The vaccine formulation comprises a
XX vaccine formulated for the prevention of a disease selected from vaccinia
XX virus (small pox), polio virus (Salk and Sabin), mumps, measles, rubella,
XX diphtheria, tetanus, Varicella-Zoster (chicken pox/shingles), pertussis
XX (whooping cough), Bacille Calmette-Guérin (BCG, tuberculosis),
XX haemophilus influenzae meningitis, rabies, cholera, Japanese
XX encephalitis virus, salmonella typhi, shigella, hepatitis A and B,
XX adenovirus, yellow fever, foot and mouth disease, herpes simplex virus,
XX respiratory syncytial virus, rotavirus, Dengue, West Nile virus, turkey
XX herpes virus (Marek's disease), influenza and/or anthrax.

XX SQ Sequence 251 AA;

Query Match 100.0%; Score 333; DB 22; Length 251;
Best Local Similarity 100.0%; Pred. No. 2.2e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 EAGLPAGKGLTGSPPGDKTGPAGQDGRPPGPPGARGQAGVMGFPKGA 59
|||||

Db 1 eaglpagkltgspgspdgktgppagdqgrppgppgargqagvmfpgpkga 59
RESULT 9
AAE02708
ID AAE02708 standard; Protein: 500 AA.
XX AC AAE02708;
XX 06-AUG-2001 (first entry)
XX Human alpha (I) type I collagen helical domain (residues 531-1030).

XX Human; recombinant gelatin; binding agent; stabilising agent; emulsifier;
XX encapsulant; film-forming agent; moisturising agent; thickening agent;
XX gelling agent; colloidal agent; adhesive agent; gel capsule; photography;
XX plasma expander; colloidal volume replacement material; graft coating;
XX medical sponge; medical plug; micro-carrier; edible composition;
XX protein supplement; fat substitute; nutritional supplement; cell culture;
XX edible coating; cosmetic; vaccine; therapy; arthritis; athrosis;
XX cartilage degeneration; joint flexibility; food industry; beverage;
XX alpha (I) type I collagen.

XX OS Homo sapiens.
XX WO200134646-A2.
XX 17-MAY-2001.

XX 10-NOV-2000; 2000WO-US30791.
XX 12-NOV-1999; 99US-0165114.
XX 15-MAY-2000; 2000US-0204437.
XX (FIBR-) FIBROGEN INC.
XX Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX WPI; 2001-329072/34.

XX Gelatin useful for pharmaceuticals, cosmetics and edible foods, is
XX prepared recombinantly -
XX Claim 21; Page 125-127; 137pp; English.
XX The patent discloses recombinant human gelatin which is useful
XX in various compositions including binding agents, encapsulants,
XX stabilising agents, film-forming agents, moisturising agents,
XX emulsifiers, thickening agents, gelling agents, colloidal agents,
XX adhesive agents, pharmaceutical compositions, hard gel capsules,
XX soft gel capsules, plasma expander, colloidal volume replacement
XX materials, graft coatings, medical sponges, medical plugs, medical
XX pharmaceutical stabilisers, micro-carriers, edible compositions,
XX protein supplements, fat substitutes, nutritional supplements,
XX edible coatings, photographic compositions, cosmetic compositions,
XX industrial composition, cell culture compositions and compositions
XX for use in the laboratory. Pharmaceutical compositions comprising
XX recombinant gelatin are used as vaccines. They are also used to
XX treat various joint conditions such as arthritis, athrosis and
XX other conditions related to the degeneration of cartilage and joint
XX flexibility. Recombinant gelatin is also used in food and beverage
XX industries. The present sequence is human alpha (I) type I collagen
XX helical domain (residues 531-1030). This sequence is a recombinant
XX gelatin.

XX SQ Sequence 500 AA;

Query Match 100.0%; Score 333; DB 22; Length 500;
Best Local Similarity 100.0%; Pred. No. 4.1e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 EAGLPAGKGLTGSPPGDKTGPAGQDGRPPGPPGARGQAGVMGFPKGA 59

```
Db 1 eaglpagkltsgpspgpdgktpgppagqdgrrpppgpaggagvmgfpgpkga 59
|||||
RESULT 10
AAB68062
ID AAB68062 standard; Protein; 500 AA.
XX
AC AAB68062;
XX
DT 09-JUL-2001 (first entry)
XX
DE Amino acid sequence of a recombinant human gelatin.
XX
KW Human; gelatin; vaccine; anaphylactic reaction.
XX
OS Homo sapiens.
XX
PN WO200134801-A2.
XX
PD 17-MAY-2001.
XX
PF 10-NOV-2000; 2000WO-US30843.
XX
PR 12-NOV-1999; 99US-0165114.
XX
PR 15-MAY-2000; 2000US-0204437.
XX
PA (FIBR-) FIBROGEN INC.
XX
PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX
DR WPI; 2001-308784/32.
XX
PT Vaccine formulations (I) comprising recombinant human gelatin, useful
PT for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies
PT and cholera, the gelatin is non-immunogenic and confers stability at
PT ambient temperatures -
XX
PS Claim 1; Page 118-120; 130pp; English.
XX
CC The present sequence represents a human recombinant gelatin polypeptide.
CC The recombinant gelatin polypeptide is used to produce vaccine
CC formulations of the invention. The recombinant human gelatin is
CC non-immunogenic (therefore reducing anaphylactic reactions) and confers
CC stability at ambient temperatures. The vaccine formulation comprises a
CC vaccine formulated for the prevention of a disease selected from vaccinia
CC virus (small pox), polio virus (Salk and Sabin), mumps, measles, rubella,
CC diphtheria, tetanus, Varicella-Zoster (chicken pox/shingles), pertussis
CC (whooping cough), Bacille Calmette-Guérin (BCG, tuberculosis),
CC haemophilus influenzae meningitis, rabies, cholera, Japanese
CC encephalitis virus, salmonella typhi, shigella, hepatitis A and B,
CC adenovirus, yellow fever, foot and mouth disease, herpes simplex virus,
CC respiratory syncytial virus, rotavirus, Dengue, West Nile virus, turkey
CC herpes virus (Marek's disease), influenza and/or anthrax.
XX
SQ Sequence 500 AA;
Query Match 100.0%; Score 333; DB 22; Length 500;
Best Local Similarity 100.0%; Pred. No. 4.1e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 EAGLPAGKLTSGPSGPDGKTGTPPGAGQDGRPGPPGARGQAGVMGFPGPKGAA 59
|||||
Db 1 eaglpagkltsgpspgpdgktpgppagqdgrrpppgpaggagvmgfpgpkga 59
|||||
RESULT 11
AAE02703
ID AAE02703 standard; Protein; 501 AA.
XX
AC AAE02703;
XX
```

```
DT 06-AUG-2001 (first entry)
XX
DE Human alpha1 (I) type I collagen helical domain (residues 179-679).
XX
KW Human; recombinant gelatin; binding agent; stabilising agent; emulsifier;
KW encapsulant; film-forming agent; moisturising agent; thickening agent;
KW gelling agent; colloidal agent; adhesive agent; gel capsule; photography;
KW plasma expander; colloidal volume replacement material; graft coating;
KW medical sponge; medical plug; micro-carrier; edible composition;
KW protein supplement; fat substitute; nutritional supplement; cell culture;
KW edible coating; cosmetic; vaccine; therapy; arthritis; atrophis;
KW cartilage degeneration; joint flexibility; food industry; beverage;
KW alpha1 (I) type I collagen.
XX
OS Homo sapiens.
XX
PN WO200134646-A2.
XX
PD 17-MAY-2001.
XX
PF 10-NOV-2000; 2000WO-US30791.
XX
PR 12-NOV-1999; 99US-0165114.
XX
PR 15-MAY-2000; 2000US-0204437.
XX
PA (FIBR-) FIBROGEN INC.
XX
PI Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
XX
DR WPI; 2001-329072/34.
XX
PT Gelatin useful for pharmaceuticals, cosmetics and edible foods, is
PT prepared recombinantly -
XX
PS Claim 21; Page 121-123; 137pp; English.
XX
CC The patent discloses recombinant human gelatin which is useful
CC in various compositions including binding agents, encapsulants,
CC stabilising agents, film-forming agents, moisturing agents,
CC emulsifiers, thickening agents, gelling agents, colloidal agents,
CC adhesive agents, pharmaceutical compositions, hard gel capsules,
CC soft gel capsules, plasma expander, colloidal volume replacement
CC materials, graft coatings, medical sponges, medical plugs,
CC pharmaceutical stabilisers, micro-carriers, edible compositions,
CC protein supplements, fat substitutes, nutritional supplements,
CC edible coatings, photographic compositions, cosmetic compositions,
CC industrial composition, cell culture compositions and compositions
CC for use in the laboratory. Pharmaceutical compositions comprising
CC recombinant gelatin are used as vaccines. They are also used to
CC treat various joint conditions such as arthritis, atrophis and
CC other conditions related to the degeneration of cartilage and joint
CC flexibility. Recombinant gelatin is also used in food and beverage
CC industries. The present sequence is human alpha1 (I) type I collagen
CC helical domain (residues 179-679). This sequence is a recombinant
CC gelatin.
XX
SQ Sequence 501 AA;
Query Match 100.0%; Score 333; DB 22; Length 501;
Best Local Similarity 100.0%; Pred. No. 4.1e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 EAGLPAGKLTSGPSGPDGKTGTPPGAGQDGRPGPPGARGQAGVMGFPGPKGAA 59
|||||
Db 353 eaglpagkltsgpspgpdgktpgppagqdgrrpppgpaggagvmgfpgpkga 411
|||||
RESULT 12
AAB68057
ID AAB68057 standard; Protein; 501 AA.
XX
AC AAB68057;
```

XX	09-JUL-2001	(first entry)
XX	DT	
XX	DE	Amino acid sequence of a recombinant human gelatin.
XX	DE	
XX	KW	Human; gelatin; vaccine; anaphylactic reaction.
XX	OS	Homo sapiens.
XX	Key	Location/Qualifiers
XX	FT	Misc-difference 85
FT	/note=	"this residue is given as unknown as it is illegible in the specification"
FT		
PX	WO200134801-A2.	
PN		
PD	17-MAY-2001.	
XX		
XX	10-NOV-2000;	2000WO-US30843.
XX	12-NOV-1999;	99US-0165114.
PR	15-MAY-2000;	2000US-0204437.
PR		
PX	{FIBR-} FIBROGEN INC.	
PA		
XX	Chang RC,	Kivirikko KI, Neff TB, Olsen DR, Polarek JW;
PI		
XX	WIPI;	2001-308784/32.
DR		
XX	Vaccine formulations (I) comprising recombinant human gelatin, useful for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies and cholera, the gelatin is non-immunogenic and confers stability at ambient temperatures .	
PT		
PT		
PT		
PT		
XX		
XX	Claim 11;	Page 114-116; 130pp; English.
PS		
CC	The present sequence represents a human recombinant gelatin polypeptide. The recombinant gelatin polypeptide is used to produce vaccine formulations of the invention. The recombinant human gelatin is non-immunogenic (therefore reducing anaphylactic reactions) and confers stability at ambient temperatures. The vaccine formulation comprises a vaccine formulated for the prevention of a disease selected from vaccinia virus (small pox), polio virus (Salk and Sabin), mumps, measles, rubella, diphtheria, tetanus, Varicella-Zoster (Chicken pox/shingles), pertussis (whooping cough), Bacille Calmette-Geurin (BCG, tuberculosis), haemophilus influenzae meningitis, rabies, cholera, Japanese encephalitis virus, salmonella typhi, shigella, hepatitis A and B, adenovirus, yellow fever, foot and mouth disease, herpes simplex virus, respiratory syncytial virus, rotavirus, Dengue, West Nile virus, turkey herpes virus (Marek's disease), influenza and/or anthrax.	
CC		
SQ	Sequence	501 AA;
	Query Match	100.0%; Score 333; DB 22; Length 501;
	Best Local Similarity	100.0%; Pred. No. 4.1e-23;
	Matches 59;	Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY	1	EAGLPGAKGLTSGSPGDGTGPDPGAGQDGRPPGGPFGARQGAGVMGFPGPKAA 59 353 eaglpagkltsgspgpgdgtgpdpaggdrppgpggagvgagvmgfpgpkga 411
Db		
RESULT 13		
AAE02718		
ID	AAE02718 standard;	Protein; 662 AA.
XX		
AC	AAE02718;	
XX		
XX	06-AUG-2001	(first entry)
DT		
XX		
DE	Human alpha (I) type I collagen helical domain (residues 531-1192).	
XX		

XX	Human; gelatin; vaccine; anaphylactic reaction.	
KW	Homo sapiens.	
XX		
OS		
XX		
PH	Key Location/Qualifiers	
FT	Misc-difference 53	
FT	/note= "this residue is given as unknown as it is illegible in the specification"	
XX		
PN	WC0200134801-A2.	
XX		
PD	17-MAY-2001.	
XX		
PF	10-NOV-2000; 2000WO-US30843.	
XX		
PR	12-NOV-1999; 99US-0165114.	
PR	15-MAY-2000; 2000US-0204437.	
XX		
PA	(FIBR-) FIBROGEN INC.	
XX		
PI	Chang RC, Kivirikko KI, Neff TB, Olsen DR, Polarek JW;	
XX		
DR	WFI; 2001-308784/32.	
XX		
PT	Vaccine formulations (I) comprising recombinant human gelatin, useful for vaccinating against e.g. mumps, measles, rubella, tetanus, rabies and cholera, the gelatin is non-immunogenic and confers stability at ambient temperatures -	
PT		
XX		
PS	Claim 11; Page 128-130; 130pp; English.	
XX		
CC	The present sequence represents a human recombinant gelatin polypeptide. The recombinant gelatin polypeptide is used to produce vaccine formulations of the invention. The recombinant human gelatin is non-immunogenic (therefore reducing anaphylactic reactions) and confers stability at ambient temperatures. The vaccine formulation comprises a vaccine formulated for the prevention of a disease selected from vaccinia virus (small pox), polio virus (Salk and Sabin), mumps, measles, rubella, diphtheria, tetanus, varicella-Zoster (chicken pox/shingles), pertussis (whooping cough), Bacille Calmette-Guérin (BCG, tuberculosis), Haemophilus influenzae meningitis, rabies, cholera, Japanese encephalitis virus, salmonella typhi, shigella, hepatitis A and B, adenovirus, yellow fever, foot and mouth disease, herpes simplex virus, respiratory syncytial virus, rotavirus, Dengue, West Nile virus, turkey herpes virus (Marek's disease), influenza and/or anthrax.	
XX		
SQ	Sequence	662 AA;
Query Match	100.0%;	Score 333; DB 22; Length 662;
Best Local Similarity	100.0%;	Pred. No. 5.3e-23;
Matches	59; Conservative	0; Mismatches 0; Indels 0; Gaps 0
Qy	1	EAGLPAGKGLTSGSPGPDGTGTGGAGDGRPPGPPGARGQAGVGMGFPQKAA 59
Db	1	eaglpgakgltspspgpdgtgtppgaqdgdrppppppgargqagvmgfpqpkkaa 59
RESULT 15		
AAy84541		
ID	AAy84541 standard; Protein; 1057 AA.	
XX		
AC	AAy84541;	
XX		
DT	25-JUL-2000 (first entry)	
XX		
DE	Amino acid sequence of a human collagen 1 (alpha1) protein.	
XX		
KW	Extracellular matrix protein; self aggregation; hydroxylated proline;	
KW	trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;	
KW	collagen; fibrinogen; fibronectin; post translational hydroxylation.	
XX		

```
XX PD 12-APR-2000.
XX PF
XX PP
XX PR 07-OCT-1999; 99EP-0119184.
XX PA
XX PA 09-OCT-1998; 98US-0169768.
XX PA (USSU ) US SURGICAL CORP.
XX PI Gruskin EA, Buechter DD, Zhang G, Connolly K;
XX WPI; 2000-259138/23.
XX DR N-PSDB; AAA12503.
XX PT Production of extracellular matrix proteins containing
XX PT 4-trans-hydroxyproline results in native self aggregating proteins,
XX PT useful on medical implants -
XX PS Example 10; Fig 39A-E; 260pp; English.
XX CC The specification describes a method for producing an extracellular
XX CC matrix protein or its fragment. The extracellular matrix protein is
XX CC capable of self aggregating in a cell which does not ordinarily
XX CC hydroxylated prolines. The method comprises optimising a nucleic acid
XX CC sequence for expression in the cell by substitution of codons preferred
XX CC by that cell for naturally occurring codons not preferred by the cell;
XX CC incorporating the nucleic acid sequence into the cell; and contacting
XX CC the cell with a hypertonic growth medium containing at least one amino
XX CC acid, selected from the group consisting of trans-4-hydroxyproline and
XX CC 3-hydroxyproline to allow at least one of the amino acids to be
XX CC assimilated into the cell and incorporated into the extracellular matrix
XX CC protein. The method may be used to make host cells assimilate and
XX CC incorporate trans-4-hydroxyproline into proteins. This is especially
XX CC useful in the recombinant production of proteins such as collagen,
XX CC fibronectin and fibronectin whose ability to self aggregate and produce
XX CC functional proteins depends on the post translational hydroxylation of
XX CC proline. The method is also useful in studying the structure and function
XX CC of polypeptides which do not normally contain trans-4-hydroxyproline.
XX CC The present sequence represents human collagen 1 (alpha1) helical region,
XX CC which may be produced using the method of the invention.
XX SQ Sequence 1057 AA;
Query Match 100.0%; Score 333; DB 21; Length 1057;
Best Local Similarity 100.0%; Pred. No. 8.2e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Qy 1 EAGLPGAKGLTGSFGSPGDKTGPAGQDGRPPGPPGARGQAGVMGFPKGA 59
Db 370 eaglpgakltgspgspgdkgtgppgagdgrrppppgargagvmgfpkga 428
RESULT 17
AA84403
ID AAY84403 standard; Protein; 1058 AA.
AC AAY84403;
XX 12-JUL-2000 (first entry)
XX DE Amino acid sequence of human type 1 (alpha1) collagen polypeptide.
XX KW Alpha1 collagen; 3,4-dehydro-L-proline; epoxidation; 3,4-epoxyproline;
XX KW collagen; mussel adhesive protein; bioadhesive.
XX OS Homo sapiens.
XX PN WO200014201-A1.
XX PD 16-MAR-2000.
XX PF 07-SEP-1999; 99WO-US20462.
```

```
XX PR 09-SEP-1998; 98US-0099652.
XX PA (USSU ) US SURGICAL CORP.
XX PA (PAOL/) PAOLELLA D N.
XX PA (GRUS/) GRUSKIN E A.
XX PA (BUEC/) BUECHTER D D.
XX PI Paolella DN, Gruskin EA, Buechter DD;
XX WPI; 2000-271051/23.
XX DR N-PSDB; AAZ99843.
XX PT Incorporating non-natural amino acid into polypeptide, useful e.g. for
XX PT production of bioadhesives, by epoxidation or substitution of
XX PT dehydroproline residues -
XX PS Disclosure; Fig 6; 66pp; English.
XX CC The present sequence represents a human type 1 (alpha1) collagen protein.
XX CC Peptides derived from the protein were used to demonstrate incorporation
XX CC of 3,4-dehydro-L-proline into the peptide, using the method of the
XX CC invention. The specification describes a method for the incorporation
XX CC of non-natural amino acid into a polypeptide. The method comprises
XX CC reacting at least one 3,4-dehydroproline residue in the polypeptide
XX CC with an epoxidation reagent from a polypeptide containing at least
XX CC one 3,4-epoxyproline residue. The method is used for studying the
XX CC effects of non-natural amino acids on structure and function of
XX CC polypeptides. The method is also useful for commercial production of
XX CC collagen or mussel adhesive proteins (which are useful as bioadhesives),
XX CC and for incorporating a wide variety of groups, including therapeutic
XX CC ligands and biological probes, into polypeptides.
XX SQ Sequence 1058 AA;
Query Match 100.0%; Score 333; DB 21; Length 1058;
Best Local Similarity 100.0%; Pred. No. 8.2e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Qy 1 EAGLPGAKGLTGSFGSPGDKTGPAGQDGRPPGPPGARGQAGVMGFPKGA 59
Db 371 eaglpgakltgspgspgdkgtgppgagdgrrppppgargagvmgfpkga 429
RESULT 18
AA89472
ID AAR89472 standard; Protein; 1107 AA.
AC AAR89472;
XX 01-OCT-1996 (first entry)
XX DE Collagen/decorin(aa46-93) fusion protein.
XX KW Transforming growth factor; TGF-beta-1; collagen IA; osteogenesis;
XX KW bone formation; tissue repair; fusion protein.
XX OS Synthetic.
XX FH Key Location/Qualifiers
XX FT Domain 1..1057
XX FT /label= Collagen-IA
XX FT /note= "collagen IA alpha-helical domain"
XX FT Peptide 1058..1059
XX FT /label= Linker_peptide
XX FT Domain 1060..1107
XX FT /label= Decorin
XX FT /note= "amino acids P46 to G93 of mature
XX FT decorin"
XX FT Misc-difference 887
XX FT /note= "unidentified amino acid"
XX FT Misc-difference 890
```



```
XX Key Location/Qualifiers
FH Domain
FT 1..1057
FT /label= Collagen-1A
FT /note= "collagen 1A alpha-helical domain"
FT Peptide
FT 1058..1059
FT /label= Linker_peptide
FT Domain
FT 1060..1171
FT /label= TGF-beta-1
FT /note= "human mature TFF-beta-1"
FT Misc-difference 887
FT /note= "unidentified amino acid"
FT Misc-difference 890
FT /note= "unidentified amino acid"
XX CA2151547-A.
PN
XX
XX 11-DEC-1995.
XX
XX 12-JUN-1995; 95CA-2151547.
XX
XX 10-JUN-1994; 94US-0259263.
XX
XX (USSU ) US SURGICAL CORP.
XX
XX Espino P, Gruskin EA;
XX WPI: 1996-140144/15.
XX N-PSDB; AAT16516.
XX
XX Chimaeric DNA encoding protein contg. extracellular matrix protein
XX domain - and cellular regulatory factor domain, partic. useful as
XX osteogenic agents, also related vectors, transformed cells and
XX chimaeric proteins.
XX Disclosure; Fig 6; 59pp; English.
XX
XX A fusion protein (AAR89470) comprises the alpha-helical region of
XX human collagen I(a) linked to the human mature transforming
XX growth factor beta-1 (TGF-beta-1). It can be expressed in
XX Escherichia coli transformants carrying a vector incorporating a
XX chimeric gene (AAT16516) coding for the fusion. The TGF-beta-
XX moiety increases efficacy of the body's normal soft tissue
XX repair response and also induces osteogenesis. The collagen
XX moiety provides an integral substructure or scaffolding for the
XX TGF and cells involved in reconstruction and growth. The fusion
XX protein provides sustained release and delivery of TGF-beta-1
XX to a target tissue.
XX Sequence 1171 AA;
XX
XX Query Match 100.0%; Score 333; DB 17; Length 1171;
XX Best Local Similarity 100.0%; Pred. No. 9.1e-23;
XX Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 EAGLPGAKGLTSGSPSPGDKTGPAGQDGRPCPGPGAGQAGVMGFPKGA 59
Db 370 eaglpgaklgtsgspgpdgkgtgppagqdgrrpppgpaggagvmgfpkpgkaa 428
RESULT 23
AAY84538
ID AAY84538 standard; Protein; 1171 AA.
XX
XX AAY84538;
XX
XX 25-JUL-2000 (first entry)
XX
XX A chimeric collagen 1 (alpha1)/TGF-beta1 protein.
XX Extracellular matrix protein; self aggregation; hydroxylated proline;
XX trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;
```

```
KW collagen; fibrinogen; fibronectin; post translational hydroxylation; ss.
KW transforming growth factor-beta1; TGF-beta1; chimera.
XX
XX Chimeric - Homo sapiens.
OS Chimeric - Unidentified.
XX
XX Key Location/Qualifiers
FT Misc-difference 858
FT /note= "Gly encoded by GCT"
XX
XX EP992586-A2.
PN
XX
XX 12-APR-2000.
PD
XX
XX 07-OCT-1999; 99EP-0119184.
XX
XX 09-OCT-1998; 98US-0169768.
XX
XX (USSU ) US SURGICAL CORP.
XX
XX Gruskin EA, Buechter DD, Zhang G, Connolly K;
XX WPI: 2000-259138/23.
XX N-PSDB; AAA12498.
XX
XX Production of extracellular matrix proteins containing
XX 4-trans-hydroxyproline results in native self aggregating proteins,
XX useful on medical implants -
XX Claim 23; Fig 15; 260pp; English.
XX
XX The specification describes a method for producing an extracellular
XX matrix protein or its fragment. The extracellular matrix protein is
XX capable of self aggregating in a cell which does not ordinarily
XX hydroxylated prolines. The method comprises optimising a nucleic acid
XX sequence for expression in the cell by substitution of codons preferred
XX by that cell for naturally occurring codons not preferred by the cell;
XX incorporating the nucleic acid sequence into the cell; and contacting
XX the cell with a hypertonic growth medium containing at least one amino
XX acid, selected from the group consisting of trans-4-hydroxyproline and
XX 3-hydroxyproline to allow at least one of the amino acids to be
XX assimilated into the cell and incorporated into the extracellular matrix
XX protein. The method may be used to make host cells assimilate and
XX incorporate trans-4-hydroxyproline into proteins. This is especially
XX useful in the recombinant production of proteins such as collagen,
XX fibrinogen and fibronectin whose ability to self aggregate and produce
XX functional proteins depends on the post translational hydroxylation of
XX proline. The method is also useful in studying the structure and function
XX of polypeptides which do not normally contain trans-4-hydroxyproline.
XX The present sequence represents chimeric collagen 1 (alpha1)/transforming
XX growth factor-beta1 (TGF-beta1) protein, which may be produced using the
XX method of the invention.
XX Sequence 1171 AA;
```

```
Query Match 100.0%; Score 333; DB 21; Length 1171;
Best Local Similarity 100.0%; Pred. No. 9.1e-23;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 EAGLPGAKGLTSGSPSPGDKTGPAGQDGRPCPGPGAGQAGVMGFPKGA 59
Db 370 eaglpgaklgtsgspgpdgkgtgppagqdgrrpppgpaggagvmgfpkpgkaa 428
RESULT 24
AAR71701
ID AAR71701 standard; protein; 1341 AA.
XX
XX AAR71701;
XX
XX 17-OCT-1995 (first entry)
XX
```


Collagen alpha 1 (I) chain precursor.

Collagen; antibody; immunoassay; metabolism; diagnosis; monitoring; disorder; osteoporosis; metastatic progression; Paget's disease; hyperthyroidism; bone; resorption; rheumatoid arthritis; osteoarthritis; vasculitis syndrome.

Homo sapiens.

Key	Location/Qualifiers
1	1.1
2	2.1
3	3.1
4	4.1
5	5.1
6	6.1
7	7.1
8	8.1
9	9.1
10	10.1
11	11.1
12	12.1
13	13.1
14	14.1
15	15.1
16	16.1
17	17.1
18	18.1
19	19.1
20	20.1
21	21.1
22	22.1
23	23.1
24	24.1
25	25.1
26	26.1
27	27.1
28	28.1
29	29.1
30	30.1
31	31.1
32	32.1
33	33.1
34	34.1
35	35.1
36	36.1
37	37.1
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87	87.1
88	88.1
89	89.1
90	90.1
91	91.1
92	92.1
93	93.1
94	94.1
95	95.1
96	96.1
97	97.1
98	98.1
99	99.1
100	100.1

```

NAME DIFFERENCE 2020
/note= "Unidentified amino acid."

```

W09508115-A.

23-MAR-1995

19-SEP-1994: 94WO-DK00348.

17-SEP-1993: 93DK-0001040

(Oste-) Osteometer AS

Bonde M. Oviist P:

WPT: 1995-131456/17

Assaying collagen fragments in body fluid by immunoassay - using antibodies raised against synthetic peptide(s) contg. potential crosslinking sites, to diagnose and monitor disorders of collagen metabolism, e.g. osteoporosis.

Disclosure (Appendix A): Page 49: 87pp: English

Determination of collagen fragments in body fluids can be achieved by immunoassay using antibodies directed against synthetic peptides derived from collagen which contain sites of potential crosslinking. The method is used to diagnose and monitor treatment of disorders of collagen metabolism (degradation of type I collagen may indicate osteoporosis, metastatic progression, Paget's disease, hyperthyroidism or other conditions involving excessive bone resorption; degradation of type II collagen may indicate rheumatoid arthritis or osteoarthritis; and of type III collagen, vasculitis syndrome). The method can also be used to assess the toxicity of a compound and to test drugs for their effect on collagen metabolism.

Sequence 1341 AA:

```

every Match      100.0%; Score 333; DB 16; Length 1341;
1st Local Similarity 100.0%; Pred. No. 1e-22;
atches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

```

1 EAGLPAAKGLTSPGSPGPDGKTGPPGAGQDGRPPGPPGARGQAGVMGFPGKGA 59
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 408 eaqlpaakqltspgsapdqktapqpaqddrppqpqpqargcavmgfgapkaaa 466

FH	Key	Location/Qualifiers
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47	47	47
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54	54	54
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56	56	56
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59	59	59
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61	61	61
62	62	62
63	63	63
64	64	64
65	65	65
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80	80	80
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82	82	82
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84	84	84
85	85	85
86	86	86
87	87	87
88	88	88
89	89	89
90	90	90
91	91	91
92	92	92
93	93	93
94	94	94
95	95	95
96	96	96
97	97	97
98	98	98
99	99	99
100	100	100

FT	misc reference 324	/note= "unidentified residue"
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E1  MISC attribute 22/      "unidentified residue"
E2  /note=

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FT      MISC difference 1127
FT      /note= "unidentified residue"

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FT	MISC	REFERENCE	1200	/note=	"unidentified residue"
----	------	-----------	------	--------	------------------------

PN US6110689-A.

PD 29-AUG-2000.

04 - NOV - 1997; 97US-0963825.

PR 21-JAN-1994; 94US-0187319.

PA (OSTE-) OSTEOMETER AS.

PI Bonde M, Qvist P;

DR WPI; 2000-586349/55.

Assaying type I collagen fragments for diagnosing osteoporosis in postmenopausal woman, involves contacting body fluid with synthetic collagen peptide and antibody and quantifying by competitive binding assay -

PS Disclosure; Column 23-37; 41pp; English.

The present sequence is that of human type I collagen alpha-1. The invention is based on the discovery of the presence of particular collagen fragments in body fluids of patients compared with those of healthy subjects. These fragments are generated upon collagen degradation and are partly characterised by the presence of potential sites for crosslinking. A method for assaying collagen fragments in a body fluid sample is based on the competitive binding to immunological binding partners of collagen fragments in the sample and of synthetic peptides derived from collagen and containing crosslinkable sites (see AA96105-11). When considering the degradation of type I collagen, the assay can be used as a means of identifying excessive bone resorption, indicating the presence of osteoporosis or the metastatic progress of a malignancy. Other conditions characterized by excessive bone resorption include paget's disease and hyperparathyroidism.

Sequence	1341 AA;
...	

Query Match	100.0%	Score 333;	DB 21;	Length 1341;
Best Local Similarity	100.0%;	Pred. No. 1e-22;		
Matches 59;	Conservative	0;	Mismatches 0;	Indels 0;
				Gaps 0;

Qy 1 EAGLPGAKGLTSGSPGPDGKTPPGPAGDGRPPGPPGARGQAGVMGFPKGAA 59
 Dd 407 eaqlpgakqltqspgspgpdgktpppgaaqdgrrpppgpaaargaaavmgfqpkgaa 465

```

XX FH Key Location/Qualifiers
XX FT 1..1057
XX FT /label= Collagen-IA
XX FT /note= "collagen IA alpha-helical domain"
XX FT Peptide
XX FT 1058..1059
XX FT /label= Linker_peptide
XX FT Domain
XX FT 1060..1388
XX FT /label= Decorin
XX FT Misc-difference 887
XX FT /note= "unidentified amino acid"
XX FT Misc-difference 890
XX FT /note= "unidentified amino acid"
XX PN CA2151547-A.
XX PN 11-DEC-1995.
XX PD 12-JUN-1995; 95CA-2151547.
XX PF 10-JUN-1994; 94US-0259263.
XX PR (USSU ) US SURGICAL CORP.
XX PA Espino P, Gruskin EA;
XX PI WPI; 1996-140144/15.
XX DR N-PSDB; AAT16517.
XX CC Chimaeric DNA encoding protein contg. extracellular matrix protein
XX PT domain - and cellular regulatory factor domain, partic. useful as
XX PT osteogenic agents, also related vectors, transformed cells and
XX PT chimaeric proteins.
XX PS Disclosure; Fig 7; 59pp; English.
XX CC A fusion protein (AAR89471) comprises the alpha-helical region of
XX CC human collagen I(a) linked to human dermatan sulphate proteoglycan
XX CC (decorin). It can be expressed in Escherichia coli transformants
XX CC carrying a vector incorporating a chimeric gene (AAT16517) coding for
XX CC the fusion. The decorin binds to type I collagen and thus affects
XX CC Elbril formation. It inhibits the cell attachment-promoting
XX CC activity of collagen and fibrinogen by binding to such molecules
XX CC near their cell binding sites. The collagen moiety provides an
XX CC integral substratum or scaffolding for the decorin. The fusion
XX CC protein acts to reduce scarring of healing tissue.
XX SQ Sequence 1388 AA;

Query Match 100.0%; Score 333; DB 17; Length 1388;
Best Local Similarity 100.0%; Pred. No. 1.1e-22;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 EAGLPKAGLTGSPGPDGKTGPPGAGQGRGPPGPPGARGQAGVMGPPGPKGAA 59
Db 370 eaglpagkltgspgpdgktgppgagqgrgppgppgargqagvmgfpgpkga 428

RESULT 27
AAY56800
ID AAY56800 standard; protein; 1411 AA.
XX AC AAY56800;
XX AC AAY56800;
XX DT 27-MAR-2000 (first entry)
XX DE Human preproalpha 1 (I) collagen.
XX KW Fibrillar collagen; C propeptide; SSAD; telopeptide; gelatin;
XX FT sequence selection and alignment domain; prosthetic implant;
XX KW foodstuff; medicine; type I collagen; human.

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OS Homo sapiens.
XX PN EP967226-A2.
XX PD 29-DEC-1999.
XX PF 04-MAY-1999; 99EP-0303470.
XX PR 08-MAY-1998; 98US-0084828.
XX PR 10-APR-1999; 99US-0289578.
XX PA (COHE-) COHESION TECHNOLOGIES INC.
XX PI Olsen DR, Hitzeman RA, Chisholm GE;
XX DR WPI; 2000-074666/07.
XX PT New method for production of fibrillar collagen, useful for preparing
XX PT telopeptide collagen fibrils and gelatin -
XX PS Example 1; Fig 3A-B; 30pp; English.
XX CC The invention provides a method for the production of fibrillar collagen.
XX CC The method comprises: (a) culturing a recombinant host cell comprising a
XX CC DNA encoding a fibrillar collagen monomer lacking a C propeptide SSAD
XX CC (sequence selection and alignment domain); and (b) producing the
XX CC fibrillar collagen. The methods are used to produce fibrillar collagen,
XX CC from which telopeptide collagen fibrils can be derived. Host cells,
XX CC comprising DNA encoding a collagen monomer lacking SSAD or N propeptide
XX CC is used to produce gelatin. Collagen is used in biological research
XX CC as a substrate for in vitro cell culture and as a component of
XX CC biocompatible materials for use in prosthetic implants, sustained drug
XX CC release matrices, artificial skin and wound dressing and healing devices.
XX CC Gelatin is particularly useful for foodstuffs and medicine, for coating
XX CC tablets and making capsules. The methods, comprising the use of collagen
XX CC monomers lacking the N and/or C propeptides, result in a large increase
XX CC in the production of type I collagen. The present sequence represents
XX CC the human preproalpha 1 (I) collagen (GenBank Accn no: AF017178).
XX SQ Sequence 1411 AA;

Query Match 100.0%; Score 333; DB 21; Length 1411;
Best Local Similarity 100.0%; Pred. No. 1.1e-22;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 EAGLPKAGLTGSPGPDGKTGPPGAGQGRGPPGPPGARGQAGVMGPPGPKGAA 59
Db 528 eaglpagkltgspgpdgktgppgagqgrgppgppgargqagvmgfpgpkga 586

RESULT 28
AAE02535
ID AAE02535 standard; Protein; 1449 AA.
XX AC AAE02535;
XX DT 10-AUG-2001 (first entry)
XX DE Porcine alpha1(I) collagen.
XX KW Porcine; alpha1(I) collagen; gelatin; cytostatic; viral infection;
XX KW pharmaceutical; food industry; cosmetic; autoimmune disorder; vaccine;
XX KW medical; arterial sealant; bone graft; dermal implant; haemostat; cancer;
XX KW rheumatoid arthritis; beverage; photographic application.
XX OS Sus scrofa.
XX FH Key Location/Qualifiers
XX FT Misc-difference 829..830
XX FT /note= "Encoded by ggcgaacctggtgatgctggtgctaaaggcgatg
XX FT ctggtccccccggccctgctgga"

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PN WO200134647-A2.
XX 17-MAY-2001.
XX 10-NOV-2000; 2000WO-US30792.
XX 12-NOV-1999; 99US-0439058.
XX 10-NOV-2000; 2000US-0439058.
XX (FIBR-) FIBROGEN INC.
XX Bell MP, Neff TB, Polarek JW, Seeley TW;
XX WPI; 2001-335911/35.
XX N-PSDB; AAD06576.
XX Novel isolated and purified bovine or porcine collagens and gelatins
XX useful in medical, pharmaceutical, food and cosmetic industries, as
XX vaccine, and for treating autoimmune disorders, infections and cancer
XX
XX Example 3; Fig 8; 168pp; English.
XX The present sequence is porcine alpha(I) collagen. The present
XX invention relates to recombinant synthesis of collagens and gelatins
XX derived from animals. Collagen is useful in medical, pharmaceutical,
XX food and cosmetic industries. Collagen is an important component of
XX arterial sealants, bone grafts, drug delivery system, dermal implants,
XX haemostats, and incontinence implants, and for treating autoimmune
XX disorders such as rheumatoid arthritis. Collagen is useful in food
XX products such as sausage casings, and in cosmetics or facial and skin
XX formulations for treating viral infections, autoimmune diseases and
XX cancer. Gelatin is useful in the manufacture or as a component of
XX various pharmaceutical and medical devices and products, in food and
XX beverage industries, in hair care and skin care products, as a glue or
XX adhesive in various manufacturing processes, as a light-sensitive coating
XX in various electronic devices, as photoresist base in photolithographic
XX processes, in printing and photographic applications, in laboratory
XX application, and as a component in various gels used for biochemical and
XX electrophoretic analysis, including enzymographic gels.
XX
XX Sequence 1449 AA;
XX
XX Query Match 100.0%; Score 333; DB 22; Length 1449;
XX Best Local Similarity 100.0%; Pred. No. 1.1e-22;
XX Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
XX
XX QY 1 EAGLPKAGLTGSPGPGDKGTGPPGAGQGRPGPPGARGQAGVMGPPGPKGAA 59
XX |||||
XX Db 533 eaglpagkltgspgpgdgtgppgagqgrpgppgargagvmgfpgpkga 591
XX |||||
XX
XX RESULT 29
XX AAE02532
XX ID AAE02532 standard; Protein; 1463 AA.
XX AC AAE02532;
XX XX
XX DT 10-AUG-2001 (first entry)
XX DE Bovine alpha(I) collagen.
XX KW Bovine: alpha(I) collagen; gelatin; cytostatic; viral infection;
XX pharmaceutical; food industry; cosmetic; autoimmune disorder; vaccine;
XX KW medical; arterial sealant; bone graft; dermal implant; haemostat; cancer;
XX KW rheumatoid arthritis; beverage; photographic application.
XX XX
XX OS Bos sp.
XX XX
XX FH Key Location/Qualifiers
XX FT Misc-difference 629
```

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FT XX /note= "Encoded by Ctr"
PN WO200134647-A2.
PD 17-MAY-2001.
XX 10-NOV-2000; 2000WO-US30792.
XX 12-NOV-1999; 99US-0439058.
XX 10-NOV-2000; 2000US-0439058.
XX (FIBR-) FIBROGEN INC.
XX Bell MP, Neff TB, Polarek JW, Seeley TW;
XX WPI; 2001-335911/35.
XX N-PSDB; AAD06573.
XX Novel isolated and purified bovine or porcine collagens and gelatins
XX useful in medical, pharmaceutical, food and cosmetic industries, as
XX vaccine, and for treating autoimmune disorders, infections and cancer
XX
XX Claim 6; Fig 2; 168pp; English.
XX The present sequence is bovine alpha(I) collagen. The present
XX invention relates to recombinant synthesis of collagens and gelatins
XX derived from animals. Collagen is useful in medical, pharmaceutical,
XX food and cosmetic industries. Collagen is an important component of
XX arterial sealants, bone grafts, drug delivery system, dermal implants,
XX haemostats, and incontinence implants, and for treating autoimmune
XX disorders such as rheumatoid arthritis. Collagen is useful in food
XX products such as sausage casings, and in cosmetics or facial and skin
XX formulations for treating viral infections, autoimmune diseases and
XX cancer. Gelatin is useful in the manufacture or as a component of
XX various pharmaceutical and medical devices and products, in food and
XX beverage industries, in hair care and skin care products, as a glue or
XX adhesive in various manufacturing processes, as a light-sensitive coating
XX in various electronic devices, as photoresist base in photolithographic
XX processes, in printing and photographic applications, in laboratory
XX application, and as a component in various gels used for biochemical and
XX electrophoretic analysis, including enzymographic gels.
XX
XX Sequence 1463 AA;
XX
XX Query Match 100.0%; Score 333; DB 22; Length 1463;
XX Best Local Similarity 100.0%; Pred. No. 1.1e-22;
XX Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
XX
XX QY 1 EAGLPKAGLTGSPGPGDKGTGPPGAGQGRPGPPGARGQAGVMGPPGPKGAA 59
XX |||||
XX Db 530 eaglpagkltgspgpgdgtgppgagqgrpgppgargagvmgfpgpkga 588
XX |||||
XX
XX RESULT 30
XX AAW68485
XX ID AAW68485 standard; Protein; 1464 AA.
XX AC AAW68485;
XX XX
XX DT 08-DEC-1998 (first entry)
XX DE Human recombinant collagen protein.
XX KW Primer: PCR; amplification; human; collagen; mammal; plant; prosthesis;
XX KW cardiac valve; ligament; tendon; skin; gingival implant; perfumes;
XX KW nerve regeneration; antibiotic; growth factor; cancer; inflammatory;
XX KW gelatin; glue; food.
XX XX
XX OS Synthetic.
XX OS Homo sapiens.
```

XX	Key	Location/Qualifiers	Matches	59;	Conservative	0;	Mismatches	0;	Indels	0;	Gaps	0;
FT	Peptide	1..22	QY	1	EAGLPKAGLTGSPGSPGDKTGTGPPGAGQDGRPGPPGARGQAGVMGPPGPKGAA	59						
FT	Protein	/note= "signal peptide"										
FT		23..999	Db	531	eaglpgaklgtsgpspgdpdgtgtgppgagqdgrrpppppggargqagvmgfpqpkga	589						
FT	Cleavage-site	161										
FT		/note= "cleavage site for aminopeptidase"										
FT	Cleavage-site	1218										
FT		/note= "cleavage site for carboxypeptidase"										
XX	WO9827202-A1.		RESULT 31									
XX			AAU14136									
XX			ID	AAU14136	standard; Protein; 1464 AA.							
XX			XX	AAU14136;								
XX	25-JUN-1998.		XX	24-OCT-2001	(first entry)							
XX	17-DEC-1997;	97WO-FR02331.	XX	Human novel protein #7.								
XX	17-DEC-1996;	96FR-0016224.	XX	Human; novel protein; Antianaemic; osteopathic; antiinflammatory;								
XX	(BIOC-) BIOCEM SA.		KW	immunomodulatory; cytostatic; neuroprotective; vulnerary; nootropic;								
XX	Bournat P, Comte J, Exposito JY, Garrone R, Gruber V;		KW	anticonvulsant; antiarthritic; cerebroprotective; antifungal; antiviral;								
XX	Merot B, Ruggiero F;		KW	antibacterial; antiallergic; dermatological; haemostatic; antiasthmatic;								
XX	WPI; 1998-362771/31.		KW	thrombolytic; immunogen; antibody; gene therapy; neurological disorder;								
XX	N-PSDB; AAV60814.		KW	Parkinson's disease; inflammatory disorder; cancer; asthma; osteoporosis;								
XX			XX	tissue regeneration; immune disorder.								
XX			OS	Homo sapiens.								
XX			XX	WO200155437-A2.								
XX			PN	02-AUG-2001.								
XX			PD	25-JAN-2001; 2001WO-US02623.								
XX			PF	25-JAN-2000; 2000US-0491404.								
XX			XX	(HYSE-) HYSEQ INC.								
XX			PR	Tang YT, Liu C, Drmanac RT;								
XX			XX	WPI; 2001-451939/48.								
XX			PI	N-PSDB; AAS22441.								
XX			DR	Isolated polypeptides useful for treating anti-inflammatory diseases,								
XX			DR	nervous system disorders, and for regenerating bone and cartilage -								
XX			XX	Example 4; Page 525-527; 894pp; English.								
XX			XX	The invention relates to polynucleotides encoding novel human								
XX			CC	proteins or their active domains. The polypeptides, polynucleotides and								
XX			CC	antibodies raised against the polypeptides are used in a method of								
XX			CC	treatment of a mammal and prevention of disorders caused by the aberrant								
XX			CC	protein expression or activity. The polypeptides can be used as								
XX			CC	molecular weight markers, food supplements, and in antibody production.								
XX			CC	The polypeptides are used to identify compounds which bind to the								
XX			CC	polypeptides. Polynucleotides of the invention are used as probes and								
XX			CC	primers, for sequencing, for chromosome or gene mapping, in the								
XX			CC	production of recombinant proteins, and in generating anti-sense DNA or								
XX			CC	RNA and in gene therapy. Polypeptides of the invention can be used to								
XX			CC	target drugs to a tumour, in assays to determine biological activity, to								
XX			CC	raise antibodies/elicit an immune response, to determine quantitative								
XX			CC	protein levels, as tissue markers, and to isolate receptors or ligands.								
XX			CC	Polypeptides of the invention may also be useful in treating platelet								
XX			CC	disorders, stem cell disorders, regenerating bone, cartilage, tendon,								
XX			CC	ligament and/or nerve tissue, wound healing, treating burns, promoting								
XX			CC	the proliferation, differentiation and survival of stem cells, as a								
XX			CC	contraceptive, treating osteoporosis and osteoarthritis, anaemia,								
XX			CC	Alzheimer's, Parkinson's and Huntington's diseases, amyotrophic lateral								
XX			CC	sclerosis, stroke, immune deficiencies resulting from bacterial, viral or								
XX			CC	fungal infection or from autoimmunity, cancer, allergy, asthma,								
XX			CC	graft-versus-host disease, eczema, haemophilia, thrombosis,								
XX			CC	anti-inflammatory diseases, nervous system disorders, and infection.								
XX			CC	The present sequence represents a protein of the invention.								

Query Match 100.0%; Score 333; DB 19; Length 1464;
Best Local Similarity 100.0%; Pred. NO. 1.le-22;

SQ Sequence 1464 AA;

Query Match 100.0%; Score 333; DB 22; Length 1464;
Best Local Similarity 100.0%; Pred. No. 1.1e-22;
Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 EAGLPGAKGLTSGSPGPDGKTGPPGAGQDGRPGPGARGAGVGMFGPKGAA 59
|||||
Db 531 eaglpgaklgtspgspgpdgktgppgagqgrpgpppgargagvmgfpgpkga 589

RESULT 32
AAB82454
ID AAB82454 standard; Protein; 1464 AA.
XX AC AAB82454;
XX DT 22-AUG-2001 (first entry)
XX DE Human pro-alpha-1 chain of type I procollagen.
XX KW COL1A1 gene; collagen; procollagen; human.
XX OS Homo sapiens.
XX FH Key Location/Qualifiers
FT Peptide 1..22
FT Protein /label= Signal_peptide
FT /label= Mature_protein
XX WO200144455-A2.
XX PN 21-JUN-2001.
XX PD 12-DEC-2000; 2000WO-GB04741.
XX PF 15-DEC-1999; 99GB-0029487.
XX PR (ASTR) ASTRAZENECA AB.
XX PA (ASTR) ASTRAZENECA UK LTD.
XX PI Ber1 R;
XX WPI; 2001-398145/42.
XX N-PSDB; AAF90491.
XX Novel antisense DNA oligonucleotide useful for inhibiting the
PT expression of wild type COL1A1 gene, for treating, reducing the risk
PT of, and preventing collagen disorders -
PS Disclosure; Page 21-26; 30pp; English.

CC The present sequence is that of the pro-alpha-1 chain of human
CC type I procollagen. The present invention relates to antisense
CC oligonucleotides (ASOs) and their use in inhibiting expression of
CC type I procollagen. The ASOs comprise 18-25 nucleotides and are
CC complementary to a specific region within the type I collagen
CC pro-alpha-1 chain gene (see AAF90491), especially those given in
CC AAF90492-503. They are capable of inhibiting the expression of
CC the pro-alpha-1 chain in a cell that expresses it. The ASOs are
CC used in a claimed method of treating, or reducing a risk of, a
CC collagen disorder. Such disorders may include those caused by
CC overproduction of collagen fibres, such as liver cirrhosis, kidney,
CC liver and heart fibrosis, scleroderma, hypertrophic scars and
CC keloids.

SQ Sequence 1464 AA;

Query Match 100.0%; Score 333; DB 22; Length 1464;
Best Local Similarity 100.0%; Pred. No. 1.1e-22;

Matches 59; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 EAGLPGAKGLTSGSPGPDGKTGPPGAGQDGRPGPGARGAGVGMFGPKGAA 59
|||||
Db 531 eaglpgaklgtspgspgpdgktgppgagqgrpgpppgargagvmgfpgpkga 589

RESULT 33
AAY84539
ID AAY84539 standard; Protein; 1388 AA.
XX AC AAY84539;
XX DT 25-JUL-2000 (first entry)
XX DE Amino acid sequence of a chimeric collagen 1 (alpha1)/decorin protein.
XX KW Extracellular matrix protein; self aggregation; hydroxylated proline;
KW trans-4-hydroxyproline; 3-hydroxyproline; recombinant protein production;
KW collagen; fibrinogen; fibronectin; post translational hydroxylation;
KW decorin; chimera.
XX OS Chimeric - Homo sapiens.
XX OS Chimeric - Unidentified.
XX FH Key Location/Qualifiers
FT Misc-difference 87 /note= "Gly encoded by GCG"
FT Misc-difference 305 /note= "Glu encoded by CAA"
FT Misc-difference 363 /note= "Gly encoded by GGT"
FT Misc-difference 378 /note= "Glu encoded by GGT"
FT Misc-difference 429 /note= "Gly encoded by CGA"
FT Misc-difference 444 /note= "Gly encoded by GCG"
FT Misc-difference 543 /note= "Gly encoded by GCC"
FT Misc-difference 546 /note= "Gly encoded by GCT"
FT Misc-difference 606 /note= "Gly encoded by GAC"
FT Misc-difference 702 /note= "Gly encoded by GCT"
FT Misc-difference 815 /note= "Pro encoded by CTT"
FT Misc-difference 858 /note= "Gly encoded by GCT"
FT Misc-difference 1066 /note= "Gly encoded by GCC"
XX EP992586-A2.
XX 12-APR-2000.
XX 07-OCT-1999; 99EP-0119184.
XX 09-OCT-1998; 98US-0169768.
XX (USSU) US SURGICAL CORP.
XX Gruskin EA, Buechter DD, Zhang G, Connolly K;
XX WPI; 2000-259138/23.
XX N-PSDB; AAA12499.
XX Production of extracellular matrix proteins containing
PT 4-trans-hydroxyproline results in native self aggregating proteins,
PT useful on medical implants -
XX Claim 25; Fig 17A-B; 260pp; English.

XX The specification describes a method for producing an extracellular
 CC matrix protein or its fragment. The extracellular matrix protein is
 CC capable of self aggregating in a cell which does not ordinarily
 CC hydroxylated prolines. The method comprises optimising a nucleic acid
 CC sequence for expression in the cell by substitution of codons preferred
 CC by that cell for naturally occurring codons not preferred by the cell;
 CC incorporating the nucleic acid sequence into the cell; and contacting
 CC the cell with a hypertonic growth medium containing at least one amino
 CC acid, selected from the group consisting of trans-4-hydroxyproline and
 CC 3-hydroxyproline to allow at least one of the amino acids to be
 CC assimilated into the cell and incorporated into the extracellular matrix
 CC protein. The method may be used to make host cells assimilate and
 CC incorporate trans-4-hydroxyproline into proteins. This is especially
 CC useful in the recombinant production of proteins such as collagen,
 CC fibronogen and fibronectin whose ability to self aggregate and produce
 CC functional proteins depends on the post translational hydroxylation of
 CC proline. The method is also useful in studying the structure and function
 CC of polypeptides which do not normally contain trans-4-hydroxyproline.
 CC The present sequence represents a chimeric collagen I (alpha1)/decorin
 CC protein, which may be produced using the method of the invention.
 XX
 SQ Sequence 1388 AA;

Query Match 97.8%; Score 325; DB 21; Length 1388;
 Best Local Similarity 98.3%; Pred. No. 5.7e-22;
 Matches 58; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

QY 1 EAGLPAGAKGLTSGSPGPGDKTGPAGODGRPGPPGARGAGVMGPPGKGA 59
 Db 370 eaglpagakeitgspgpgpdkgtgpppagqdrpppgppgargagvmgfpkga 428

RESULT 34

AAAY06239
 ID AAY06239 standard; Protein; 595 AA.

AC AAY06239;

XX 23-AUG-1999 (first entry)

XX Mouse recombinant type I collagen COL1A1-2.

KW Type I collagen; COL1A2-1; mouse; silver halide; emulsion;
 peptide; photography.

OS Mus sp.

FH Key Location/Qualifiers

FT Cleavage-site 38..41

FT /note= "MGPR protease recognition sequence"

FT Cleavage-site 122..125

FT /note= "MGPR protease recognition sequence"

FT EP926543-A1.

XX 30-JUN-1999.

XX 15-DEC-1998; 98EP-0204263.

XX 24-DEC-1997; 97NL-1007908.

XX (FUJF) FUJI PHOTO FILM BV.

XX Bouwstra JB, De Wolf FA, Mooibroek A, Van Den Bosch TJ;

PI Van Heerde GV, Van Rijn AC, Werten MWT, Wind RD;

XX WPI; 1999-349297/30.

XX New tabular silver halide emulsion, useful for production of

PT components for photographic products

XX

PS Claim 9; Fig 10; 30pp; English.

XX This is the amino acid sequence of recombinant mouse type I
 CC collagen COL1A1-2, obtained by expression of COL1A1-2 cDNA from
 CC vector pCOL1A1-2 in transformed Pichia pastoris GS115 host cells.
 CC The invention relates to a new tabular silver halide emulsion
 CC comprising silver halide grains nucleated in the presence of a
 CC nucleation peptizer and grown in the presence of a growth peptizer,
 CC at least one of the peptizers being a pure collagen-like material,
 CC such as the present protein, prepared by genetic engineering of a
 CC native collagen-encoding nucleic acid. Also claimed is production
 CC of the recombinant collagen-like polypeptide comprising expression
 CC of a collagen-like polypeptide nucleic acid sequence by a
 CC microorganism selected from Hansenula, Trichoderma, Aspergillus and
 CC preferably P. pastoris, the collagen-like polypeptide being obtained
 CC at a level greater than 0.95 g/l (especially over 3 g/l) and free of
 CC helix structure. The emulsion is suitable for photographic
 CC application. Recombinant DNA technology enables the efficient
 CC production of large amounts of substantially pure collagen material,
 CC providing a high level of expression without requiring expensive
 CC media, expression hosts or non-secreting expression hosts. The
 CC collagen can be selected and/or adapted for optimal use in each
 CC particular stage of the production process of the photographic
 CC product. Removal of collagen MGPR motifs that are recognised by a
 CC P. pastoris protease will also increase expression levels.
 XX

SQ Sequence 595 AA;

Query Match 96.4%; Score 321; DB 20; Length 595;
 Best Local Similarity 96.6%; Pred. No. 6e-22;
 Matches 57; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 1 EAGLPAGAKGLTSGSPGPGDKTGPAGODGRPGPPGARGAGVMGPPGKGA 59
 Db 352 eaglpagakeitgspgpgpdkgtgpppagqdrpppgppgargagvmgfpkgt 410

RESULT 35

AAAY06240

ID AAY06240 standard; Protein; 822 AA.

XX AAY06240;

XX 23-AUG-1999 (first entry)

XX Mouse recombinant type I collagen COL1A1-3.

KW Type I collagen; COL1A2-3; mouse; silver halide; emulsion;
 peptide; photography.

OS Mus sp.

FH Key Location/Qualifiers

FT Cleavage-site 38..41

FT /note= "MGPR protease recognition sequence"

FT Cleavage-site 122..125

FT /note= "MGPR protease recognition sequence"

FT EP926543-A1.

XX 30-JUN-1999.

XX 15-DEC-1998; 98EP-0204263.

XX 24-DEC-1997; 97NL-1007908.

XX (FUJF) FUJI PHOTO FILM BV.

XX Bouwstra JB, De Wolf FA, Mooibroek A, Van Den Bosch TJ;

PI Van Heerde GV, Van Rijn AC, Werten MWT, Wind RD;

XX WPI; 1999-349297/30.

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